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<p>(21) International Application Number: PCT/US97/01384</p> <p>(22) International Filing Date: 28 January 1997 (28.01.97)</p> <p>(30) Priority Data: 08/592,900 29 January 1996 (29.01.96) US </p> <p>(71) Applicant: ARIZONA BOARD OF REGENTS, on behalf of UNIVERSITY OF ARIZONA [US/US]; 888 North Euclid Avenue, Main Gate 515, P.O. Box 210158, Tucson, AZ 85721-0158 (US).</p> <p>(72) Inventors: DELLA PENNA, Dean; 4135 Longknife Road, Reno, NV 89509 (US). NORRIS, Susan, R.; Apartment #2012, 555 East Limberlost Road, Tucson, AZ 85705 (US).</p> <p>(74) Agent: BERSON, Bennett, J.; Quarles & Brady, P.O. Box 2113, Madison, WI 53701-2113 (US).</p>		<p>(81) Designated States: AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, UZ, VN, ARIPO patent (KE, LS, MW, SD, SZ, UG), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG).</p> <p>Published <i>With international search report.</i></p>	
<p>(54) Title: CLONED PLANT P-HYDROXYPHENYL PYRUVIC ACID DIOXYGENASE</p> <p>(57) Abstract</p> <p>A cDNA clone from <i>Arabidopsis thaliana</i>, pHPP1.5, SEQ ID NO:1, which encodes the enzyme p-hydroxyphenyl pyruvic acid dioxygenase, is disclosed. A vector and microbial host containing a DNA sequence coding for the expression of <i>Arabidopsis thaliana</i> p-hydroxyphenyl pyruvic acid dioxygenase, and a genetic construct containing a DNA sequence coding for the expression of <i>Arabidopsis thaliana</i> p-hydroxyphenyl pyruvic acid dioxygenase, together with a promoter located 5' to the DNA coding sequence and a 3' termination sequence, are also disclosed. A method of creating a transgenic plant in which production of plastoquinones, vitamin E, and carotenoids has been modified, is also disclosed.</p>			

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5 CLONED PLANT P-HYDROXYPHENYL PYRUVIC ACID DIOXYGENASE

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Field Of The Invention

The present invention relates to a molecular approach for modifying the synthesis of vitamin E, plastoquinone, and carotenoids in plants by use of a full-length cloned cDNA which encodes a p-hydroxyphenyl pyruvic acid dioxygenase enzyme.

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Background Of The Invention

The chloroplasts of higher plants contain many unique, interconnected biochemical pathways that produce an array of secondary metabolite compounds which not only perform vital functions within the plant but are also important from agricultural and nutritional perspectives. Three such secondary metabolites are the lipid soluble, chloroplastically synthesized compounds vitamin E (α -tocopherol or α -toc), plastoquinones (PQ), and carotenoids, which together perform many crucial biochemical functions in the chloroplast. PQ and vitamin E are quinone compounds synthesized by a common pathway in the plastid; carotenoids are tetraterpenoids synthesized by a separate plastid-localized pathway.

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Plastoquinone (PQ) often accounts for up to 50% of the total plastidic quinone pool in green tissues. The primary function of PQ is as a fundamental component of the photosynthetic electron transport chain, acting as an electron carrier between

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photosystem II and the cytochrome b₆f complex. PQ likely has other less well studied functions in plastids, namely in acting as a direct or intermediate electron carrier for a variety of other biosynthetic reactions in the chloroplast.

Vitamin E is the second major class of chloroplastic quinones, accounting for up to 40% of the quinone pool in plastids. The essential nutritional value of tocopherols was recognized around 10 1925, and the compound responsible for Vitamin E activity was first identified as α-tocopherol in 1936. α-Toc has a well-documented role in mammals as an antioxidant, and a similar, though less well understood antioxidant role in plants. Liebler, et 15 al., Toxicology 23:147-169, 1993; Hess, Anti-oxidants in Higher Plants, CRC Press: 111-134, 1993.

Carotenoids are a separate, diverse group of lipophilic pigments synthesized in plants, fungi, and bacteria. In photosynthetic tissues, carotenoids function as accessory pigments in light harvesting and play important roles in photo-protection by quenching free radicals, singlet oxygen, and other reactive species. Siefermann-Harms, Physiol. Plantarum. 69:561-568, 1987. In the plastids of non-photosynthetic tissues, high levels of carotenoids often accumulate providing the intense orange, yellow, and red coloration of many fruits, vegetables, and flowers (Pfander, Methods in Enzym., 213A, 3-13, 1992). In addition to their many functions in plants, 20 carotenoids and their metabolites also have important functions in animals, where they serve as the major source of Vitamin A (retinol), and have been identified as providing protection from some forms of cancer due to their antioxidant activities. Vitamin 25 E's antioxidant activities are also thought to protect against some forms of cancer, and may act synergistically with carotenoids in this regard.

Liebler, et al., Toxicology 23:147-169, 1993; Krinsky, J. Nutr. 119:123-126, 1989.

Tocopherol and Plastoquinone Synthesis

5 α -Tocopherol and plastoquinone are the most abundant quinones in the plastid and are synthesized by the common pathway shown in Figure 1. The precursor molecule for both compounds, homogentisic acid (HGA), is produced in the chloroplast from the shikimic acid pathway intermediate p-hydroxyphenyl pyruvic acid (pOHPP), in an oxidation/decarboxylation reaction catalyzed by the enzyme p-hydroxyphenyl pyruvic acid dioxygenase (pOHPP dioxygenase). Homogentisic acid is subject to
10 phytylation/prenylation (phytylpyrophosphate and solanylpyrophosphate, C₂₀ and C₄₅, respectively) coupled to a simultaneous decarboxylation by a phytyl/prenyl transferase to form the first true tocopherol and plastoquinone intermediates,
15 2-demethylphytylplastoquinol and 2-demethylplastoquinol-9, respectively. A single ring methylation occurs on 2-demethylplastoquinol to yield plastoquinol-9 (PQH₂) which is then oxidized to plastoquinone-9 (PQ). This oxidation is reversible and is the basis of electron transport by
20 plastoquinone in the chloroplast.
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The preferred route, as established in spinach, for α -tocopherol formation from 2-demethylphytylplastoquinol appears to be 1) ring methylation of the intermediate, 2- α -demethylphytylplastoquinol, to yield phytylplastoquinol, 2) cyclization to yield d-tocopherol and, finally, 3) a second ring methylation to yield α -tocopherol. Ring methylation in both tocopherol and plastoquinone synthesis is carried out by a single enzyme that is specific for the site of methylation on the ring, but has
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relatively broad substrate specificity and accommodates both classes of quinone compounds. This methylation enzyme is the only enzyme of the pathway that has been purified from plants to date.

5 d'Harlingue, et al., J.Biol.Chem. 26:15200, 1985. All enzymatic activities of the α -toc/PQ pathway have been localized to the inner chloroplast envelope by cell fractionation studies except for pOHPP dioxygenase and the tocopherol cyclase enzyme. Difficulties with cell
10 fractionation methods, low activities for some of the enzymes, substrate stability and availability and assay problems, make studying the pathway biochemically difficult.

Vitamin E and PQ levels, ratios, and total
15 amounts vary by orders of magnitude in different plants, tissues and developmental stages. Such variations indicate that the vitamin E and PQ pathway is both highly regulated and has the potential for manipulation to modify the absolute levels and ratios
20 of the two end products. The pathway in Figure 1 makes it clear that production of homogentisic acid by pOHPP dioxygenase is likely to be a key regulatory point for bulk flow through the pathway, both because HGA production is the first committed step in α -toc/PQ
25 synthesis, and also because the reaction is essentially irreversible. Therefore modifying the levels of HGA by modifying pOHPP dioxygenase activity should have a direct impact on the total α -toc/PQ biosynthetic accumulation in plant tissues, and, as
30 described below, because of the connection of PQ and carotenoid synthesis, should also affect carotenoid synthesis in plant tissues.

Carotenoid Biosynthesis; Quinones as Electron Carriers

In plants, carotenoids are synthesized and
35 accumulate exclusively in plastids via the pathway shown on the left-hand side of Figure 1. The first

committed step in carotenoid synthesis is the condensation of two molecules of the C₂₀ hydrocarbon geranylgeranyl pyrophosphate (GGDP) by the enzyme phytoene synthase, to form the colorless C₄₀ hydrocarbon, phytoene. In oxygenic photosynthetic organisms (e.g. plants, algae, and cyanobacteria), phytoene undergoes two sequential desaturation reactions, catalyzed by phytoene desaturase, to produce ζ -carotene through the intermediate phytofluene. Subsequently, ζ -carotene undergoes two further desaturations, catalyzed by ζ -carotene desaturase, to yield the red pigment lycopene. Lycopene is cyclized to produce either α -carotene or β -carotene, both of which are subject to various hydroxylation and epoxidation reactions to yield the carotenoids and xanthophylls most abundant in photosynthetic tissues of plants, lutein, β -carotene, violaxanthin and neoxanthin.

The genes encoding the first two enzymes of the carotenoid pathway (phytoene synthase and phytoene desaturase) have been isolated and studied from a number of plant and bacterial sources in recent years. Sandmann, Eur. J. Biochem. 223:7-24, 1994. Phytoene desaturase has been the most intensively studied, both because it is a target for numerous commercially important herbicides, and also because the phytoene desaturation reaction is thought to be a rate limiting step in carotenoid synthesis. Molecular and biochemical studies suggest that two types of phytoene desaturase enzymes have evolved by independent evolution: the crtI-type found in anoxygenic photosynthetic organisms (e.g. Rhodobacter and Erwinia), and the pds-type found in oxygenic photosynthetic organisms. Despite their differences in primary amino acid sequence, all phytoene desaturase enzymes contain a dinucleotide binding domain (FAD or NAD/NADP), which in Capsicum annuum has

been shown to be FAD. Hugueney et al., Eur. J. Biochem. 209:399-407, 1992. Presumably, the bound dinucleotide in both types of phytoene desaturase enzymes is reduced during desaturation and reoxidized by an unknown reductant present in the plastid or bacterium.

Several lines of evidence have suggested a role for quinones in the phytoene desaturation reaction in higher plants. Using isolated daffodil chromoplasts, Mayer and co-workers demonstrated that in an anaerobic environment, oxidized artificial quinones were required for the desaturation of phytoene while reduced quinones were ineffective. Mayer et al., Eur. J. Biochem. 191:359-363, 1990. Further supporting evidence comes from studies with the triketone class of herbicides (e.g. Sulcotrione), which cause phytoene accumulation in treated tissues but unlike the well-studied pyridazone class (e.g. Norflorazon (NFZ)) do not directly affect the phytoene desaturase enzyme. Rather, triketone herbicides competitively inhibit pOHPP dioxygenase, an enzyme common to the synthesis of both plastoquinone and tocopherols, suggesting that one or more classes of quinones may play a role in carotenoid desaturation reactions. Schulz et al., FEBS 318:162-166, 1993; Secor, Plant Physiol. 106: 1429-1433; Beyer et al., IUPAC Pure and Applied Chemistry 66:1047-1056, 1994.

Despite the well-studied, wide-spread importance of vitamin E, plastoquinone, and carotenoids to human nutrition, agriculture, and biochemical processes within plant cells, much remains unclear about their biosynthesis and accumulation in plant tissues. This uncertainty has in turn limited the potential for manipulation of the synthesis and levels of these important compounds in plants.

Summary of the Invention

In one embodiment, this invention provides a biologically pure sample of DNA which DNA comprises a sequence coding for the expression of *Arabidopsis thaliana* p-hydroxyphenyl pyruvic acid dioxygenase.

5 In other embodiments, this invention provides a vector and microbial host containing a DNA sequence sufficiently homologous to SEQ ID NO:1 so as to code for the expression of *Arabidopsis thaliana* p-hydroxyphenyl pyruvic acid dioxygenase, and a genetic construct containing a DNA sequence sufficiently homologous to SEQ ID NO:1 so as to code for the expression of *Arabidopsis thaliana* p-hydroxyphenyl pyruvic acid dioxygenase, together with a promoter located 5' to the DNA coding sequence and a 3' termination sequence.

10 In another embodiment, this invention provides a method of creating a transgenic plant in which the levels of the pOHPP dioxygenase enzyme are elevated sufficient such that production of plastoquinones, vitamin E, and carotenoids are modified.

15 It is an object of the present invention to genetically engineer higher plants to modify the production of plastoquinones, vitamin E, and carotenoids.

20 It is another object of the invention to provide transgenic plants that would express elevated levels of the pOHPP dioxygenase enzyme which would have resultant elevated resistance to the triketone class of herbicides (i.e. sulcotrione).

25 It is another object of the present invention to provide a method for the preparation of the enzyme p-hydroxyphenyl pyruvic acid dioxygenase (pOHPP dioxygenase), an enzyme which can be used to identify new pOHPPdioxygenase-inhibiting herbicides.

30 Other features and advantages of the invention will be apparent from the following description of the

preferred embodiments thereof and from the claims.

Brief Description of the Drawings

Fig. 1 is a diagram of the pathways for synthesis of carotenoids, vitamin E (tocopherol), and
5 plastoquinone..

Fig. 2 is a diagram of the interconnections of the pathways illustrated in Fig. 1.

Fig. 3A-3E are graphs of pigment analyses of wild-type, NFZ-wt, and pds1 tissues.

10 Fig. 4 is a physical map of the pds1 mutation relative to visible markers.

Figs. 5A-5C present the results of C18 HPLC separation of lipid soluble pigments from wild-type plants on MS2 media, homozygous pds1 mutants on MS2
15 media supplemented with pOHPP, and homozygous pds1 mutants on MS2 media supplemented with homogentistic acid (HGA).

Figs. 6A-6B present the results of C8 HPLC analyses of quinones in NFZ-wt and pds1 tissues.

20 Detailed Description Of The Invention

As described above, both Vitamin E, plastoquinones and carotenoids are synthesized and accumulated in plastids by the pathways shown in Figure 1. This specification describes the identification, isolation, characterization and functional analysis of a higher plant pOHPP dioxygenase cDNA, its role in α -toc, PQ and carotenoid synthesis, and the use of this cDNA to modify pOHPP dioxygenase activity in plant tissues and hence the accumulation of one or more of the compounds plastoquinones, vitamin E, and carotenoids in plant tissues. The overexpression of pOHPP dioxygenase in transgenic plants will modify the enzyme-to-inhibitor ratio of plant tissues exposed to triketone herbicides, as compared to non-transgenic plants,

resulting in increased herbicide resistance. The present specification also describes a genetic construct for use in the production of pOHPP dioxygenase, an enzyme useful in identifying new pOHPP dioxygenase-inhibiting herbicides.

By genetic analysis the present inventors have shown that the vitamin E, plastoquinone, and carotenoid biosynthetic pathways are interconnected and share common elements as shown in Figure 2. From 10 mutational studies in *Arabidopsis thaliana*, the present inventors identified one genetic locus, designated *pds1* (*pds*= phytoene desaturation), the disruption of which results in accumulation of the first carotenoid of the carotenoid biosynthetic pathway, phytoene. Surprisingly, though this mutation disrupts carotenoid synthesis and was originally 15 identified on this basis, it does not map to the locus encoding the phytoene desaturase enzyme. Evidence indicates that *pds1* defines a second gene product in addition to the phytoene desaturase enzyme, necessary 20 for phytoene desaturation and hence carotenoid synthesis in higher plants. This gene product proved to be pOHPP dioxygenase.

To provide a molecular mechanism for manipulating 25 synthesis and accumulation of the compounds plastoquinone, vitamin E, and carotenoids, the present inventors used a molecular genetic approach, taking advantage of the model plant system *Arabidopsis thaliana* to define, isolate and study genes required 30 for synthesis of the compounds in plants. The flowering plant *Arabidopsis thaliana* has come into wide use as a model system to explore the molecular biology and genetics of plants. *Arabidopsis* offers many advantages for genetic analysis: it can be selfed 35 and very large numbers of progeny can be obtained (up to 10,000 seeds from a single plant). Furthermore, *Arabidopsis* has a short generation time of five to six

weeks, so crosses can be set up and the progeny analyzed within reasonable periods of time. Mutation screens have identified thousands of mutations affecting many aspects of basic plant biology, including morphogenesis, photosynthesis, fertility, starch and lipid metabolism, mineral nutrition, and so on. In addition, its haploid genome is only about 10^8 base pairs.

An important aspect of the successful approach used here is that essential components were first functionally defined genetically, prior to their isolation, analysis and molecular manipulation. Briefly, potential mutants were identified by a combination of phenotypic and biochemical screening, characterized at the genetic and molecular levels, loci of interest selected, and the corresponding genes then cloned and studied further. By this approach, the inventors genetically defined and isolated cDNAs for one gene, *pds1*, whose mutation disrupts synthesis of all three classes of compounds in the plastid, tocopherols, plastoquinones and carotenoids. Based on biochemical analysis of the *pds1* mutant, the *pds1* gene was identified as affecting the activity of pOHPP dioxygenase, a crucial enzyme of the plastidic quinone pathway in plants (Figure 1), that is directly required for the synthesis of plastoquinone and α -tocopherol and indirectly for carotenoid synthesis. In particular, the deduced function of the *pds1* mutant and pOHPP dioxygenase enzyme are noted in Figure 2.

The present inventors demonstrated by biochemical complementation that the *pds1* mutation affects the enzyme p-hydroxyphenyl pyruvic acid dioxygenase (pOHPP dioxygenase), because *pds1* plants can be rescued by growth on the product but not the substrate of this enzyme, homogentisic acid (HGA) and p-hydroxyphenylpyruvate (pOHPP), respectively. pOHPP dioxygenase is the key branch point enzyme and

committed step in the synthesis of both Vitamin E and plastoquinones and several independent lines of biochemical evidence confirm *pds1* affects this enzyme (Figures 1, 5, 6). These results provide the first 5 genetic evidence that plastoquinones are essential components for carotenoid synthesis in higher plants, most likely as an electron carrier/redox element in the desaturation reaction (Figure 2). The *Arabidopsis* pOHPP dioxygenase gene/cDNA thus provides a basis for 10 modifying the production of plastoquinones, α -tocopherol and carotenoids in all higher plants.

Specifically, the specification describes the 15 genetic identification of the *Arabidopsis* pOHPP dioxygenase gene by mutational analysis, the physical isolation and functional confirmation of an *Arabidopsis* pOHPP dioxygenase cDNA, its nucleotide sequence and its use to isolate pOHPP dioxygenase genes and cDNAs from other plant species. Also included in the specification is a description of the 20 use of the *Arabidopsis* pOHPP dioxygenase cDNA, and related cDNAs from other plants, to positively or negatively modify the expression/activity of pOHPP dioxygenase by recombinant techniques (overexpression, cosuppression, antisense, etc.) in any and all plant 25 tissues, especially leaf and fruit tissues, to positively or negatively affect the production of α -toc, PQ and carotenoids.

Elevating pOHPP dioxygenase protein levels 30 increases the amount of homogentisic acid (HGA) synthesized in plant tissues. Because HGA is the limiting precursor molecule for α -toc and PQ synthesis (the end products of the pathway), increasing HGA synthesis increases the levels of α -toc (Vitamin E) and PQ in plant tissues. The increase in PQ 35 indirectly increases the synthesis of carotenoids, which require PQ for their synthesis. In addition, the increase in PQ increases photosynthetic efficiency

by increasing electron flow between photosystem II and photosystem I, because PQ is the primary electron transporter between the two photosystems. The increase in α -toc, a well-studied antioxidant in mammals, increases the ability of plants to withstand oxidative stresses, such as that caused by high light, high temperature, water stress, ozone stress, UV stress or other abiotic or biotic stresses. Elevating the levels of pOHPP dioxygenase will modify the dose response curve of herbicides targeting pOHPP dioxygenase, thus increasing the relative resistance to such herbicides in transgenic plants as compared to native plants of the same species. Inhibiting the expression of pOHPP dioxygenase is expected to have the opposite effect.

Genetic Construct

To express pOHPP dioxygenase in a plant, it is required that a DNA sequence containing the pOHPP dioxygenase coding sequence be combined with regulatory sequences capable of expressing the coding sequence in a plant. A number of effective plant promoters, both constitutive and developmentally or tissue specific, are known to those of skill in the art. A transcriptional termination sequence (polyadenylation sequence) may also be added. Plant expression vectors, or plasmids constructed for expression of inserted coding sequences in plants, are widely used in the art to assemble chimeric plant expression constructs including the coding sequence, and to conveniently transfer the constructs into plants. A sequence which codes for pOHPP dioxygenase includes, for example, SEQ ID NO:1, or versions of the designated sequence sufficient to effect coding for the expression of pOHPP dioxygenase. Commonly used methods of molecular biology well-known to those of skill in the art may be used to manipulate the DNA

sequences.

By "genetic construct" we mean any of a variety of ways of combining the protein-encoding sequences with a promoter sequence (and termination sequence, if necessary) in a manner that operably connects the promoter sequence (and termination sequence, if present) with the protein-encoding sequences.

Typically, the promoter sequence will be "upstream" of a protein-encoding sequence, while the termination sequence, if used, will be "downstream" of the protein-encoding sequences.

The protein-encoding, promoter and termination sequences may be combined on a plasmid or viral vector, and inserted into a microbial host. Other functional sequences may be added to the gene construct. Alternatively, the protein-encoding, promoter, and termination sequence, if added, may be combined with any other needed functional sequences and used without a vector.

The DNA sequence described by SEQ ID NO:1 is sufficient to effect coding for the expression of pOHPP dioxygenase. However, it is envisioned that the above sequence could be truncated and still confer the same properties. It is not known at present which specific deletions would be successful, but it is likely that some deletions to the protein would still result in effective enzymatic activity. One skilled in the art of molecular biology would be able to take the designated sequence and perform deletional analysis experiments to determine what portions of the designated sequence are essential to effect coding for the expression of pOHPP dioxygenase. One could create a genetic construct with the candidate deletion mutations and perform experiments as described below in the Examples, to test whether such deletion mutation sequences effect coding for the enzyme. Expression of the enzyme activity indicates a

successful deletion mutant or mutants. In this manner, one could determine which parts of the designated sequence is essential for expression of the enzyme.

5 It is also known that the genetic code is degenerate, meaning that more than one codon, or set of three nucleotides, codes for each amino acid. Thus it is possible to alter the DNA coding sequence to a protein, such as the sequence for pOHPP dioxygenase described here, without altering the sequence of the protein produced. Selection of codon usage may affect expression level in a particular host. Such changes in codon usage are also contemplated here.

10 15 It is further contemplated that using the *Arabidopsis* pOHPP gene coding sequence described here, that the homologous pOHPP dioxygenase sequences from other higher plants can be readily recovered. Oligonucleotides can be made from the sequence set forth below to either hybridize against cDNA or 20 genomic libraries or used for PCR amplification of homologous pOHPP dioxygenase sequences from other plants.

Once a pOHPP gene is in hand, whether from 25 *Arabidopsis* or from some other plant species, it then becomes possible to insert a chimeric plant expression genetic construct into any plant species of interest. Suitable plant transformation methods exist to insert 30 such genetic constructs into most, if not all, commercially important plant species. Presently known methods include *Agrobacterium*-mediated transformation, coated-particle gene delivery (Biolistics) and electroporation, in which an electric voltage is used to facilitate gene insertion. All these methods, and others, can insert the genetic construct into the 35 genome of the resulting transgenic plant in such a way that the genetic construct becomes an inheritable trait, transmitted to progeny of the original

transgenic plant by the normal rules of Mendelian inheritance. Thus, once a genetic construct expressing a pOHPP gene is inserted into a plant, it can become a part of a plant breeding program for transfer into any desired genetic background.

To over-express pOHPP dioxygenase, a genetic construct may be used with a higher strength promoter. To inhibit expression of endogenous pOHPP dioxygenase, an antisense genetic construct can be made, as is known by those of skill in the art, to reduce the level of pOHPP dioxygenase present in the plant tissues.

EXAMPLES

Isolation of *pds1*, a mutant defective in carotenoid synthesis

To further understand carotenoid biosynthesis and its integration with other pathways in the chloroplast in higher plants, the present inventors studied the pathway by isolating *Arabidopsis thaliana* mutants that are blocked in carotenoid synthesis.

Plants homozygous for defects in the early stages of carotenoid synthesis (e.g. prior to production of β -carotene) are lethal when grown in soil and the isolation of such mutations requires the design of screening procedures to identify plants heterozygous for soil lethal mutations. The present inventors found that most soil lethal, homozygous pigment-deficient *Arabidopsis* mutants can be grown to near maturity in tissue culture on Murashige and Skoog basal media (Murashige and Skoog, *Physiol. Plant.* 15:473-497, 1962) supplemented with sucrose (MS2 media). Under these conditions, photosynthesis and chloroplast development are essentially dispensable and all the energy and nutritional needs of the plant are supplied by the media.

Greater than 500 lines from the 10,000 member

Feldmann T-DNA tagged *Arabidopsis thaliana* population (Forsthoefel et al., Aust. J. Plant Physiol. 19:353-366, 1992) were selected for pigment analysis based on their segregation for lethal pigment mutations. Seed 5 from plants heterozygous for lethal pigment mutations were surface sterilized, grown on MS2 media, the segregating pigment mutants identified, tissue harvested from individual plants, and HPLC pigment analysis performed. Although numerous mutant lines 10 with severe pigment deficiencies were identified, only two were found to be carotenoid biosynthetic mutants. One mutant line isolated from this group, *pds1*, is described in detail here.

The hallmark phenotype for disruption of a 15 biosynthetic pathway is the accumulation of an intermediate compound prior to the site of blockage. Such blockage of the carotenoid pathway can be mimicked chemically by treatment of wild-type plants with the herbicide NFZ, an inhibitor of the phytoene 20 desaturase enzyme (Figure 1) which has been reported to cause accumulation of phytoene in treated tissues. Britton, Z. Naturforsch 34c:979-985, 1979. Figs. 3A-3E present the results of pigment analysis of wild-type, NFZ-wt, and *pds1* tissues. Abbreviations in 25 Figs. 3A-3E are as follows: N, neoxanthin; V, violaxanthin; L, lutein; Cb, chlorophyll b; Ca, chlorophyll a; β , β -carotene.

Figure 3A shows C₁₈ Reverse Phase HPLC analysis of 30 the carotenoids that accumulate in wild-type *Arabidopsis thaliana* leaves. In comparison, Figure 3B shows the pigment profile for NFZ treated wild-type (NFZ-Wt). Spectral analysis of the strongly absorbing 296nm peak at 33 minutes in NFZ-Wt tissue shows absorbance maxima at 276, 286, and 298nm, indicative 35 of phytoene (Figure 3D). Figure 3C shows pigment analysis of tissue culture grown homozygous *pds1* mutant plants. The low absorbance at 440nm in Figures

3B and C demonstrates that like NFZ-Wt, *pds1* mutants lack all chlorophylls and carotenoids that normally accumulate in wild-type tissue (compare to Figure 3A). However, unlike wild-type, *pds1* mutants contain a peak
5 with a retention time at approximately 33 minutes that absorbs strongly at 296nm. The retention time and absorbance of the 33-minute peak in the *pds1* mutant corresponds to the phytoene peak in pigment extracts of NFZ-Wt tissue (Figure 3B). Spectral analysis of
10 the 33-minute peak from *pds1* is shown in Figure 3E and is virtually identical to the spectra of phytoene from NFZ-Wt tissue (Figure 3D) as well as to the published spectra for phytoene. These results confirm the
15 chemical identity of the accumulating compound in *pds1* as phytoene and conclusively demonstrate that the *pds1* mutation disrupts carotenoid biosynthesis.

Carotenoid Analysis

For quantitative and qualitative carotenoid analysis, plant tissue is placed in a microfuge tube and ground with a micropesle in 200 μ l of 80% acetone. 20 120 μ l of ethyl acetate is added and the mixture vortexed. 140 μ l of water is added and the mixture centrifuged for 5 minutes. The carotenoid containing upper phase is then transferred to a fresh tube and vacuum dried in a Jouan RC1010 Centrifugal Evaporator.
25 The dried extract is resuspended in ethyl acetate at a concentration of 0.5mg fresh weight of tissue per μ l and either analyzed immediately by HPLC or stored at -80°C under nitrogen.

Carotenoids were separated by reverse-phase HPLC 30 analysis on a Spherisorb ODS2 5 micron C₁₈ column, 25 cm in length (Phase Separations Limited, Norwalk, CT) using a 45 minute gradient of Ethyl Acetate (0-100%) in Acetonitrile/water/triethylamine (9:1:0.01 v/v), at 35 a flow rate of 1 ml per minute (Goodwin and Britton, 1988). Carotenoids were identified by retention time

relative to known standards with detection at both 296nm and 440nm. When needed, absorption spectra for individual peaks were obtained with a Hewlett Packard 1040A photodiode array detector and compared with published spectra or available standards.

5 Quinone analysis

Quinones were extracted from tissue using a method modified from that described in Bligh et al., Can. J. Biochem. Physiol. 37:911-917, 1959. Frozen 10 plant tissue was ground in a mortar with 3 volumes of chloroform and 6 volumes of methanol and transferred to a test tube. Water and additional chloroform were added until a biphasic mixture was obtained. The quinone containing chloroform phase was then 15 collected. To increase yields, the aqueous phase was back-extracted with chloroform, the two chloroform phases pooled, and then filtered through Whatman #3 filter paper. The resulting filtrate was dried under a constant stream of nitrogen. Once dried, the pellet 20 was resuspended in methanol at a concentration of 10mg fresh weight per ml and immediately analyzed by HPLC. Quinones were resolved by reversed-phase HPLC analysis on a LiChrosorb RP-8, 5 micron column, 25cm in length, (Alltech, San Jose, CA) using an isocratic solvent of 25 10% H₂O in Methanol for the first 14 minutes, at which time the solvent was switched to 100% methanol for the remainder of the run (modified from the method described in Lichtenthaler, Handbook of Chromatography, CRC Press, 115-159, 1984). The flow 30 rate was 1ml per minute for the duration. Peaks were identified based upon the retention time of known standards with detection at 280nm for α -tocopherol and 35 260nm for plastoquinone and ubiquinone as well as by absorption spectra from a Hewlett Packard 1040A photodiode array detector. When needed, fractions represented by individual chromatographic peaks were

collected, and submitted to the Southwest Environmental Health Science Center, Analytical Core laboratory for mass spectral analysis. Results were obtained using a TSQ7000 tandem mass spectrometer (Finnigan Corp., San Jose, CA) equipped with an atmospheric pressure chemical ionization source operated in the positive ion mode. The instrument was set to unit resolution and the samples were introduced into the source in a 0.3 ml/minute methanol stream and ionized using a 5kV discharge.

Genetic analysis of *pds1*

The genetic nature of the *pds1* mutation was determined by analyzing seeds resulting from selfing *pds1* heterozygous plants. Prior to desiccation, F1 seeds were scored as either green (wild-type or heterozygous) or white (homozygous). A 3:1 segregation ratio was observed (146 green seeds: 48 white seeds), indicating that *pds1* is inherited as single recessive nuclear mutations ($\chi^2=0.01$, $p > 0.90$). Because *pds1* mutants are inhibited in the desaturation of phytoene, the inventors believed that it might be a mutation in the phytoene desaturase enzyme, which had previously been mapped to chromosome 4, between *ag* and *bp*. Wetzel et al., Plant J. 6:161-175, 1994. To test this hypothesis, the *pds1* mutation was mapped relative to visible markers. The *pds1* mutation was found to map to chromosome 1, approximately 7 cM from *dis1* toward *clv2*. Franzmann et al., Plant J. 7:341-350, 1995. These data points are summarized in Figure 4 and establish that *pds1* does not map to the phytoene desaturase enzyme locus, thus proving that the *pds1* mutation is not in the phytoene desaturase enzyme. This data provided important insight for characterization of the *pds1* mutant.

Homozygous *pds1* mutants can be rescued by Homogentisic Acid, an intermediate in plastoquinone and tocopherol biosynthesis

As described earlier, previous research suggesting a role for quinones and pOHPP dioxygenase in phytoene desaturation lead the present inventors to investigate the quinone biosynthetic pathway in the *pds1* mutant. The early stages of plastoquinone/tocopherol synthesis were functionally analyzed by growth in the presence of two intermediate compounds in the pathway, p-hydroxyphenylpyruvate (pOHPP) and homogentisic acid (HGA) (refer to Figures 1 and 2). Albino *pds1* homozygous plants were first germinated on MS2 media and then transferred to MS2 media supplemented with 100 μ M of either pOHPP or HGA. *pds1* plants remained albino when transferred to media containing pOHPP but greening occurred when *pds1* plants were transferred to media containing HGA. Figs. 5A-5C present the results of complementation of the *pds1* mutation with homogentisic acid. Each profile represents pigments extracted from 10mg fresh weight of tissue. Abbreviations used in Figs. 5A-5C are as described in Figs. 3A-3E. HPLC analysis with detection at 440nm of the carotenoids extracted from *pds1* plants grown on pOHPP and HGA are shown in Figures 5B and C, respectively. The pigment profiles of *pds1* mutants grown on pOHPP are similar to the profiles of *pds1* plants grown on MS2 media shown in Figure 3B. Comparison of the pigment profiles for *pds1* + HGA tissue and wild-type tissue (Figures 5A and 5C) indicates that growth in the presence of HGA is able to qualitatively restore a wild-type carotenoid profile to albino, homozygous *pds1* plants. These results indicate that the *pds1* mutation affects the enzyme pOHPP dioxygenase, because *pds1* mutants are not altered by growth on the substrate of this enzyme, pOHPP, but rather, are restored qualitatively to wild-type pigmentation by growth on the product of

this enzyme, HGA (refer to Figures 1 and 2). The complementation of *pds1* with HGA also indicates that intermediates or end products of this pathway (plastoquinone and/or tocopherols, refer to Figures 1 and 2) are necessary components for phytoene desaturation in plants and confirms the observation of Schultz et al. in FEBS where inhibitors of POHPP dioxygenase were shown to cause accumulation of phytoene.

HPLC analysis conclusively demonstrates that *pds1* is a mutation in the plastoquinone/ tocopherol biosynthetic pathway that also affects carotenoid synthesis

In addition to biochemical complementation of *pds1* mutants, the plastoquinone/tocopherol pathway was also directly analyzed in *pds1* tissue by utilizing C₈ HPLC to resolve total lipid extracts and identify three separate classes of quinones: ubiquinone, plastoquinone, and α -tocopherol (Vitamin E) (Figures 5 and 6). Ubiquinone and plastoquinone perform analogous electron transport functions in the mitochondria and chloroplast, respectively, but are synthesized by different pathways in separate subcellular compartments (Goodwin et al., Introduction to Plant Biochemistry, Oxford, Pergamon Press, 1983), making ubiquinone an ideal internal control in these analyses. Figure 6 shows the C₈ HPLC analysis of lipid soluble extracts from NFZ-Wt tissue and *pds1* tissue. In NFZ-Wt tissue (Figure 6A), peaks 3 and 4 were identified as ubiquinone and plastoquinone, respectively, based on retention time (26 and 27 minutes), optical spectra, and mass spectra (results not shown). NFZ-Wt tissue contained a peak (1) with a retention time of 13.5 minutes which was identified as α -tocopherol based upon the retention time of a standard. However, optical spectroscopy and mass spectrometry demonstrated that peak 1 was composed of two major components: α -tocopherol (la) and an

unidentified compound (1b). The mass of α -tocopherol was determined to be 430 as indicated by the presence of the 431 protonated molecule while the molecular mass of the unidentified compound was 412, as indicated by the presence of the 413 protonated molecule (data not shown), clearly demonstrating the presence of two compound in peak 1. This quinone analysis demonstrates that the herbicide NFZ, which specifically inhibits the phytoene desaturase enzyme, does not affect synthesis of homogentisate derived quinones. *pds1* tissue (Figure 6B) contain ubiquinone (peak 3) but lack plastoquinone (peak 4). Additionally, though *pds1* contains a peak at 13.5 minutes, optical spectroscopy and mass spectrometry data demonstrate that this peak lacks α -tocopherol (1a) and is composed solely of the compound 1b (data not shown). Therefore, homozygous *pds1* plants accumulate ubiquinone but lack both plastoquinone and α -tocopherol. This is consistent with the *pds1* mutation affecting pOHPP dioxygenase (refer to Figures 1 and 2), as suggested by the rescue of the mutation by HGA, and provide additional evidence that the *pds1* mutation disrupts pOHPP dioxygenase.

**Isolation of a truncated, putative pOHPP dioxygenase
Arabidopsis cDNA**

The observation of Schultz et al. demonstrating that inhibitors of pOHPP dioxygenase activity disrupt carotenoid synthesis and cause accumulation of phytoene provided important insight for the characterization of the *pds1* mutant which in turn provided the present inventors with important insight for the isolation of a putative cDNA for the *pds1* locus. In animals, genetic defects which inhibit the activity of pOHPP dioxygenase lead to tyrosinemia type I, a fatal inherited disease in aromatic amino acid catabolism characterized by the presence of high levels of pOHPP in the urine.

In an effort to further understand the nature of this disease, pOHPP dioxygenase cDNAs have been cloned from several mammalian and bacterial sources (summarized in Ruetschi et al., Eur. J. Biochem. 205:459-466, 1992). Amino acid identity between various mammalian pOHPP dioxygenase enzymes is >80%; in comparison, their identity to bacterial homologs is very low, less than 28%. By using mammalian and bacterial sequences to search the Expressed Sequence Tags (ESTs) computer DNA database (Newman et al., Plant Physiol. 106:1241-1255, 1994), one partial length *Arabidopsis* EST was identified and used as a probe. The partial length *Arabidopsis* probe corresponds to base pairs 1072 through 1500 of SEQ ID NO:1.

This cDNA contained only 99 amino acids of the carboxyl terminal portion of the protein coding region. The deduced protein sequence of this putative *Arabidopsis* pOHPP dioxygenase cDNA shows similar homology (~50% identity) to both the mammalian and bacterial pOHPP dioxygenases. Interestingly, the partial *Arabidopsis* sequence also contains a 15 amino acid insertion not found in the human or bacterial enzymes. Finally, alignment of six pOHPP dioxygenase sequences from mammals and bacteria identified three regions of high conservation, the highest being a 16 amino acid region near the carboxy end of pOHPP dioxygenases that shows 62.5% identity across all phyla. Ruetschi et al., Eur. J. Biochem. 205:459-466, 1992. This region is also present in the truncated *Arabidopsis* sequence. The lines of evidence suggest that the partial length *Arabidopsis* cDNA described above encodes a pOHPP dioxygenase, most likely the *pds1* locus.

**Isolation and Characterization of a full length
Arabidopsis pOHPP dioxygenase cDNA**

Utilizing the partial length *Arabidopsis* cDNA probe, an *Arabidopsis* cDNA library was screened by 5 nucleic acid hybridization for full length cDNAs. A large number of hybridizing cDNAs were isolated, and one of the longest, pHPP1.5, containing a 1,520 bp insertion, was sequenced completely; the insert is presented as SEQ ID NO: 1. pHPP1.5 encodes a 446 10 amino acid protein (presented as SEQ ID NO:2), which is slightly larger in size than mammalian and bacterial pOHPP dioxygenases. pHPP1.5 shows 34-40% identity at the amino acid level to pOHPP dioxygenases from various mammals and bacteria. In comparing four 15 bacterial pOHPP dioxygenases and one mammalian pOHPP dioxygenases (pig) which ranged in size from 346-404 amino acids, Denoya et al. identified 69 amino acids that were conserved between all five pOHPP dioxygenases. Denoya et al., J. Bacteriol. 176:5312- 20 5319, 1994. The pHPP1.5 coding region contains 52 of these 69 conserved amino acids.

Demonstration that pHPP1.5 encodes an active pOHPP dioxygenase protein and complements the *pds1* mutation

In order to definitively demonstrate that pHPP1.5 25 is the gene product encoded by the *pds1* locus and that it encodes a functional pOHPP dioxygenase protein, the pHPP1.5 cDNA was cloned into a plant transformation vector for molecular complementation experiments with the *pds1*. The full length wild-type pOHPP dioxygenase 30 cDNA will be subcloned into a plant transformation vector driven by the Cauliflower Mosaic Virus 35S (CaMV) promoter and containing all necessary termination cassettes and selectable markers (Kan^r). The CaMV promoter is a strong constitutive promoter. 35 This single construct and the vector without the pHPP1.5 insert (as a control) will be used in vacuum infiltration transformation which uses whole soil

grown plants and will be done on plants that are heterozygous for the *pds1* mutation. Bouchez et al., CR Acad. Sci. Paris, Sciences de la vie 316, 1993.

In the standard procedure, 20-30 soil grown
5 plants will be independently transformed and analyzed separately. In this case homozygous plants containing the *pds1* mutation would be lethal while heterozygous plants containing the *pds1* mutation would be segregating 2:1 for the *pds1* mutation in their
10 siliques. The inventors will use a similar number of wild type plants in a parallel transformation as a control. After transformation of the *pds1* segregating plant population, as the plants are setting seed the inventors can easily identify those heterozygous for
15 the *pds1* mutation in retrospect by inspection of their siliques which would contain green:white embryos in a 3:1 ratio.

Seed harvested from individually transformed heterozygous *pds1* plants will be germinated on
20 kanamycin and resistant seedlings transferred to soil. Segregation analysis of seed from these primary transformants (T2 seed) and T3 seed for segregation of the *pds1* phenotype (albino and phytoene accumulating) and the T-DNA encoded kanamycin resistance marker
25 (wild type pOHPP dioxygenase cDNA) will conclusively demonstrate complementation of the *pds1* mutation with the pOHPP dioxygenase cDNA. To provide additional proof that pHPP1.5 is encoded by the *pds1* locus, the pHPP1.5 cDNA has been mapped relative to the *pds1*
30 locus using recombinant inbred lines, as described in Lister et al., Plant J. 4:745-750, 1993. The pHPP1.5 cDNA mapped to the region of chromosome 1 containing the *pds1* mutation (Figure 4). Finally, the pHPP1.5 cDNA will be overexpressed in E. coli and the activity
35 of the protein determined.

Modification of pOHPP Expression

From the genetic and biochemical studies described above it is clear that only one pOHPP dioxygenase gene product is involved in chloroplastic quinone synthesis, that the *pds1* mutation defines this gene, that the pHPP1.5 cDNA is the product encoded by the *pds1* locus and that disruption of its function completely eliminates Vitamin E production and plastoquinone and carotenoid synthesis in plant tissues. Modification of pOHPP dioxygenase expression in plants by molecular techniques using pHPP1.5 can therefore be used to positively or negatively affect the production of tocopherols, plastoquinones directly and carotenoids indirectly (refer to Figures 1 and 2). Specifically, overexpression of the pOHPP dioxygenase enzyme will result in increased levels of one or more of these compounds in the tissues of transgenic plants. Alternatively, using antisense techniques, it is possible to lower the level of enzyme activity to decrease the levels of these compounds in plants. Additionally, overexpression of the pOHPP dioxygenase will enable a transgenic plant to withstand elevated levels of herbicides that target this enzyme, providing agronomically significant herbicide resistance relative to normal plants.

Two different plant systems, *Arabidopsis* and tomato, are being used to demonstrate the effects of modified pOHPP dioxygenase in plant tissues. Constitutive overexpression of pOHPP dioxygenase will be done in both plant systems utilizing the CaMV 35S promoter and the pHPP1.5 cDNA. The consequences of this altered expression on tocopherol, plastoquinone and carotenoid levels and profiles in various plant tissues will be determined as described below. In tomato, tissue specific overexpression of pOHPP dioxygenase (pHPP1.5) will be driven by the fruit specific promoter derived from the tomato β subunit

gene, which is expressed specifically in developing, but not ripening tomato fruit. This will determine the potential for modifying the levels of tocopherol, plastoquinone and carotenoids specifically in

5 developing and ripening fruit for nutritional purposes without affecting their production in other plant tissues. These combined experiments will determine whether pOHPP dioxygenase is a rate limiting step in chloroplastic homogentisic acid derived quinone

10 synthesis and the potential for manipulating chloroplastic homogentisic acid derived quinones (tocopherols and plastoquinones) and compounds that require quinones for their synthesis (carotenoids, etc) by increasing pOHPP dioxygenase activity.

15 Multiple independent transformants will be produced for each construct and plant species used. The integration and gene copy number of each chimeric gene in each line will be confirmed by southern analysis, the level of pOHPP mRNA determined by

20 Northern blot analysis, pOHPP dioxygenase activity determined as described in Schulz et al., FEBS 318:162-166, 1993, and the effects on individual chloroplastic components of interest analyzed (tocopherols, plastoquinones and carotenoids). In

25 green tissue containing constitutively expressing constructs this analysis can occur relatively soon after transformants are put into soil. Analysis of fruit specific construct lines will require much more time for fruit set to occur. Analysis of tocopherols,

30 plastoquinones and carotenoids will be by a combination of HPLC, optical and mass spectra as described in Norris et al. (1995, in press). Analysis of tocopherol levels is performed by HPLC and when needed by GC:mass spectroscopy in selected ion mode.

35 In MS analysis the absolute level of tocopherol will be quantified by isotopic dilution with a known, "heavy carbon" tocopherol standard added at the start

of the extraction. Determination based on fresh weight of tissue can also be performed. Plastoquinone levels will be quantified by C8 HPLC and optical spectra as described in Norris et al. (1995, in press). Total carotenoid levels are determined spectrophotometrically and the levels of individual carotenes quantified by C18 HPLC and optical spectra quantified to standards. In the course of these experiments we will identify high expressing lines with simple insertions that segregate as single genetic loci in progeny. This will facilitate analysis of the inheritance of the gene and phenotype in future generations.

Overexpression of pOHPP for in vitro Herbicide Analysis

pOHPP dioxygenase will be overexpressed in E. coli or other prokaryotic or eukaryotic protein production systems and purified in large amounts for use in enzymatic assays for identifying new herbicide compounds (pOHPP inhibitors) and optimizing existing chemistries through detailed kinetic analysis.

SEQUENCE LISTING

(1) GENERAL INFORMATION:

(i) APPLICANT: DellaPenna, Dean
Norris, Susan

5 (ii) TITLE OF INVENTION: Cloned Plant P-Hydroxyphenyl
Pyruvic Acid Dioxygenase

(iii) NUMBER OF SEQUENCES: 2

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15 (F) ZIP: 53701-2113

(v) COMPUTER READABLE FORM:

(A) MEDIUM TYPE: Floppy disk
(B) COMPUTER: IBM PC compatible
(C) OPERATING SYSTEM: PC-DOS/MS-DOS
20 (D) SOFTWARE: PatentIn Release #1.0, Version #1.30

(vi) CURRENT APPLICATION DATA:

(A) APPLICATION NUMBER:
(B) FILING DATE:
(C) CLASSIFICATION:

25 (viii) ATTORNEY/AGENT INFORMATION:

(A) NAME: Seay, Nicholas J
(B) REGISTRATION NUMBER: 27,386
(C) REFERENCE/DOCKET NUMBER: 920214.90158

(ix) TELECOMMUNICATION INFORMATION:

30 (A) TELEPHONE: 608-251-5000
(B) TELEFAX: 608-251-9166

(2) INFORMATION FOR SEQ ID NO:1:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 1519 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
35 (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(vi) ORIGINAL SOURCE:

40 (A) ORGANISM: Arabidopsis thaliana

(vii) IMMEDIATE SOURCE:

(B) CLONE: pHPP1.5

(ix) FEATURE:

(A) NAME/KEY: CDS
45 (B) LOCATION: 37..1374

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

CCACGCGTCC GAGTTTAGC AGAGTTGGTG AAATCA ATG GGC CAC CAA AAC GCC
 Met Gly His Gln Asn Ala 1 5 54

5 GCC GTT TCA GAG AAT CAA AAC CAT GAT GAC GGC GCT GCG TCG TCG CCG
 Ala Val Ser Glu Asn Gln Asn His Asp Asp Gly Ala Ala Ser Ser Pro 10 15 20 102

10 GGA TTC AAG CTC GTC GGA TTT TCC AAG TTC GTA AGA AAG AAT CCA AAG
 Gly Phe Lys Leu Val Gly Phe Ser Lys Phe Val Arg Lys Asn Pro Lys 25 30 35 150

15 TCT GAT AAA TTC AAG GTT AAG CGC TTC CAT CAC ATC GAG TTC TGG TGC
 Ser Asp Lys Phe Lys Val Lys Arg Phe His His Ile Glu Phe Trp Cys 40 45 50 198

20 GGC GAC GCA ACC AAC GTC GCT CGT CGC TTC TCC TGG GGT CTG GGG ATG
 Gly Asp Ala Thr Asn Val Ala Arg Arg Phe Ser Trp Gly Leu Gly Met 55 60 65 70 246

25 AGA TTC TCC GCC AAA TCC GAT CTT TCC ACC GGA AAC ATG GTT CAC GCC
 Arg Phe Ser Ala Lys Ser Asp Leu Ser Thr Gly Asn Met Val His Ala 75 80 85 294

30 TCT TAC CTA CTC ACC TCC GGT GAC CTC CGA TTC CTT TTC ACT GCT CCT
 Ser Tyr Leu Leu Thr Ser Gly Asp Leu Arg Phe Leu Phe Thr Ala Pro 90 95 100 342

35 TAC TCT CCG TCT CTC TCC GCC GGA GAG ATT AAA CCG ACA ACC ACA GCT
 Tyr Ser Pro Ser Leu Ser Ala Gly Glu Ile Lys Pro Thr Thr Thr Ala 105 110 115 390

40 TCT ATC CCA AGT TTC GAT CAC GGC TCT TGT CGT TCC TTC TTC TCT TCA
 Ser Ile Pro Ser Phe Asp His Gly Ser Cys Arg Ser Phe Phe Ser Ser 120 125 130 438

45 CAT GGT CTC GGT GTT AGA GCC GTT GCG ATT GAA GTA GAA GAC GCA GAG
 His Gly Leu Gly Val Arg Ala Val Ala Ile Glu Val Glu Asp Ala Glu 135 140 145 150 486

50 TCA GCT TTC TCC ATC AGT GTA GCT AAT GGC GCT ATT CCT TCG TCG CCT
 Ser Ala Phe Ser Ile Ser Val Ala Asn Gly Ala Ile Pro Ser Ser Pro 155 160 165 534

55 CCT ATC GTC CTC AAT GAA GCA GTT ACG ATC GCT GAG GTT AAA CTA TAC
 Pro Ile Val Leu Asn Glu Ala Val Thr Ile Ala Glu Val Lys Leu Tyr 170 175 180 582

60 GGC GAT GTT GTT CTC CGA TAT GTT AGT TAC AAA GCA GAA GAT ACC GAA
 Gly Asp Val Val Leu Arg Tyr Val Ser Tyr Lys Ala Glu Asp Thr Glu 185 190 195 630

65 AAA TCC GAA TTC TTG CCA GGG TTC GAG CGT GTA GAG GAT GCG TCG TCG
 Lys Ser Glu Phe Leu Pro Gly Phe Glu Arg Val Glu Asp Ala Ser Ser 200 205 210 678

70 TTC CCA TTG GAT TAT GGT ATC CGG CGG CTT GAC CAC GCC GTG GGA AAC
 Phe Pro Leu Asp Tyr Gly Ile Arg Arg Leu Asp His Ala Val Gly Asn 215 220 225 230 726

75 GTT CCT GAG CTT GGT CCG GCT TTA ACT TAT GTA GCG GGG TTC ACT GGT
 Val Pro Glu Leu Gly Pro Ala Leu Thr Tyr Val Ala Gly Phe Thr Gly 235 240 245 774

	TTT CAC CAA TTC GCA GAG TTC ACA GCA GAC GAC GTT GGA ACC GCC GAG Phe His Gln Phe Ala Glu Phe Thr Ala Asp Asp Val Gly Thr Ala Glu 250 255 260	822
5	AGC GGT TTA AAT TCA GCG GTC CTG GCT AGC AAT GAT GAA ATG GTT CTT Ser Gly Leu Asn Ser Ala Val Leu Ala Ser Asn Asp Glu Met Val Leu 265 270 275	870
	CTA CCG ATT AAC GAG CCA GTG CAC GGA ACA AAG AGG AAG AGT CAG ATT Leu Pro Ile Asn Glu Pro Val His Gly Thr Lys Arg Lys Ser Gln Ile 280 285 290	918
10	CAG ACG TAT TTG GAA CAT AAC GAA GGC GCA GGG CTA CAA CAT CTG GCT Gln Thr Tyr Leu Glu His Asn Glu Gly Ala Gly Leu Gln His Leu Ala 295 300 305 310	966
15	CTG ATG AGT GAA GAC ATA TTC AGG ACC CTG AGA GAG ATG AGG AAG AGG Leu Met Ser Glu Asp Ile Phe Arg Thr Leu Arg Glu Met Arg Lys Arg 315 320 325	1014
	AGC AGT ATT GGA GGA TTC GAC TTC ATG CCT TCT CCT CCG CCT ACT TAC Ser Ser Ile Gly Gly Phe Asp Phe Met Pro Ser Pro Pro Pro Thr Tyr 330 335 340	1062
20	TAC CAG AAT CTC AAG AAA CGG GTC GGC GAC GTG CTC AGC GAT GAT CAG Tyr Gln Asn Leu Lys Arg Val Gly Asp Val Leu Ser Asp Asp Gln 345 350 355	1110
	ATC AAG GAG TGT GAG GAA TTA GGG ATT CTT GTA GAC AGA GAT GAT CAA Ile Lys Glu Cys Glu Glu Leu Gly Ile Leu Val Asp Arg Asp Asp Gln 360 365 370	1158
25	GGG ACG TTG CTT CAA ATC TTC ACA AAA CCA CTA GGT GAC AGG CCG ACG Gly Thr Leu Leu Gln Ile Phe Thr Lys Pro Leu Gly Asp Arg Pro Thr 375 380 385 390	1206
30	ATA TTT ATA GAG ATA ATC CAG AGA GTA GGA TGC ATG ATG AAA GAT GAG Ile Phe Ile Glu Ile Gln Arg Val Gly Cys Met Met Lys Asp Glu 395 400 405	1254
	GAA GGG AAG GCT TAC CAG AGT GGA GGA TGT GGT GGT TTT GGC AAA GGC Glu Gly Lys Ala Tyr Gln Ser Gly Gly Cys Gly Phe Gly Lys Gly 410 415 420	1302
35	AAT TTC TCT GAG CTC TTC AAG TCC ATT GAA GAA TAC GAA AAG ACT CTT Asn Phe Ser Glu Leu Phe Lys Ser Ile Glu Glu Tyr Glu Lys Thr Leu 425 430 435	1350
	GAA GCC AAA CAG TTA GTG GGA TGA ACAAGAAGAA GAACCAACTA AAGGATTGTG Glu Ala Lys Gln Leu Val Gly *	1404
40	440 445	
	TAATTAATGT AAAACTGTTT TATCTTATCA AAACAATGTT ATACAACATC TCATTTAAAAA ACGAGATCAA TCAAAAAATA CAATCTTAAA TTCAAAACC A AAAAAAAA AAAAAA	1464 1519

(2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 446 amino acids
 (B) TYPE: amino acid
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Met Gly His Gln Asn Ala Ala Val Ser Glu Asn Gln Asn His Asp Asp
 1 5 10 15

Gly Ala Ala Ser Ser Pro Gly Phe Lys Leu Val Gly Phe Ser Lys Phe
 5 20 25 30

Val Arg Lys Asn Pro Lys Ser Asp Lys Phe Lys Val Lys Arg Phe His
 35 40 45

His Ile Glu Phe Trp Cys Gly Asp Ala Thr Asn Val Ala Arg Arg Phe
 50 55 60

Ser Trp Gly Leu Gly Met Arg Phe Ser Ala Lys Ser Asp Leu Ser Thr
 10 65 70 75 80

Gly Asn Met Val His Ala Ser Tyr Leu Leu Thr Ser Gly Asp Leu Arg
 85 90 95

Phe Leu Phe Thr Ala Pro Tyr Ser Pro Ser Leu Ser Ala Gly Glu Ile
 15 100 105 110

Lys Pro Thr Thr Ala Ser Ile Pro Ser Phe Asp His Gly Ser Cys
 115 120 125

Arg Ser Phe Phe Ser Ser His Gly Leu Gly Val Arg Ala Val Ala Ile
 130 135 140

Glu Val Glu Asp Ala Glu Ser Ala Phe Ser Ile Ser Val Ala Asn Gly
 20 145 150 155 160

Ala Ile Pro Ser Ser Pro Pro Ile Val Leu Asn Glu Ala Val Thr Ile
 165 170 175

Ala Glu Val Lys Leu Tyr Gly Asp Val Val Leu Arg Tyr Val Ser Tyr
 25 180 185 190

Lys Ala Glu Asp Thr Glu Lys Ser Glu Phe Leu Pro Gly Phe Glu Arg
 195 200 205

Val Glu Asp Ala Ser Ser Phe Pro Leu Asp Tyr Gly Ile Arg Arg Leu
 210 215 220

Asp His Ala Val Gly Asn Val Pro Glu Leu Gly Pro Ala Leu Thr Tyr
 30 225 230 235 240

Val Ala Gly Phe Thr Gly Phe His Gln Phe Ala Glu Phe Thr Ala Asp
 245 250 255

Asp Val Gly Thr Ala Glu Ser Gly Leu Asn Ser Ala Val Leu Ala Ser
 35 260 265 270

Asn Asp Glu Met Val Leu Leu Pro Ile Asn Glu Pro Val His Gly Thr
 275 280 285

Lys Arg Lys Ser Gln Ile Gln Thr Tyr Leu Glu His Asn Glu Gly Ala
 290 295 300

Gly Leu Gln His Leu Ala Leu Met Ser Glu Asp Ile Phe Arg Thr Leu
 40 305 310 315 320

Arg Glu Met Arg Lys Arg Ser Ser Ile Gly Gly Phe Asp Phe Met Pro
 325 330 335

Ser Pro Pro Pro Thr Tyr Tyr Gln Asn Leu Lys Lys Arg Val Gly Asp
 45 340 345 350

Val Leu Ser Asp Asp Gln Ile Lys Glu Cys Glu Glu Leu Gly Ile Leu
355 360 365

Val Asp Arg Asp Asp Gln Gly Thr Leu Leu Gln Ile Phe Thr Lys Pro
370 375 380

5 Leu Gly Asp Arg Pro Thr Ile Phe Ile Glu Ile Ile Gln Arg Val Gly
385 390 395 400

Cys Met Met Lys Asp Glu Glu Gly Lys Ala Tyr Gln Ser Gly Gly Cys
405 410 415

10 Gly Gly Phe Gly Lys Gly Asn Phe Ser Glu Leu Phe Lys Ser Ile Glu
420 425 430

Glu Tyr Glu Lys Thr Leu Glu Ala Lys Gln Leu Val Gly *
435 440 445

5

CLAIMS

I claim:

1. A biologically pure sample of DNA, the DNA comprising a DNA sequence coding for the expression of a plant p-hydroxyphenyl pyruvic acid dioxygenase.

10 2. A vector containing the DNA sequence of
claim 1.

3. A microbial host transformed by the vector
of claim 2.

15 4. The DNA of claim 1, wherein the p-
hydroxyphenyl pyruvic acid dioxygenase is from
Arabidopsis thaliana.

5. A transgenic tomato plant transformed with a
DNA construct including the DNA of claim 1.

20 6. A transgenic Arabidopsis plant transformed
with a DNA construct including the DNA of claim 1

7. The biologically pure DNA of claim 1 wherein
the DNA is SEQ ID NO:2.

8. A DNA plant gene expression construct
comprising:

25 a. a DNA sequence coding for the
expression of a plant p-hydroxyphenyl pyruvic acid
dioxygenase;

b. a promoter effective in plant cells
located 5' to the DNA coding sequence; and

30 c. a 3' termination sequence effective in
plant cells.

9. A DNA construct comprising:

a. a promoter capable of expressing a downstream coding sequence in a tomato plant;

5 b. a DNA sequence coding for the expression of a p-hydroxyphenyl pyruvic acid dioxygenase of plant origin; and

c. a 3' termination sequence, the construction capable of expressing a p-hydroxyphenyl pyruvic acid dioxygenase gene when transformed into 10 tomato plants.

10. A bacteria containing the construction of Claim 9.

11. A tomato plant cell containing the construction of Claim 9.

15 12. An Arabidopsis plant cell containing the construction of Claim 9.

13. A transgenic tomato plant comprising in its genome a foreign genetic construction comprising, 5' to 3', a promoter effective in tomato, a DNA coding 20 region encoding p-hydroxyphenyl pyruvic acid dioxygenase, and a transcriptional terminator, the genetic construction effective in vivo in tomato plants to stimulate expression of p-hydroxyphenyl pyruvic acid dioxygenase.

25 14. Seed of the tomato plant of claim 13.

15. Fruit of the tomato plant of claim 13.

16. The transgenic tomato plant of claim 11 wherein the DNA coding region encoding p-hydroxyphenyl pyruvic acid dioxygenase is that set forth in SEQ ID 30 NO:1.

17. A method of suppressing the production of vitamin E and plastoquinones in a plant, the method comprising the steps of:

- a. isolating a DNA sequence encoding a p-hydroxyphenyl pyruvic acid dioxygenase of plant origin;
- b. creating a genetic construction including, 5' to 3', a promoter effective in the plant's cells, a coding sequence, and a transcriptional terminator, the coding region being derived from the DNA sequence, wherein the DNA sequence from step (a) has been altered so that expression of p-hydroxyphenyl pyruvic acid dioxygenase is suppressed; and
- c. transforming a cell of the plant with the genetic construction, whereby the plant's cell produces lowered levels of vitamin E and plastoquinones.

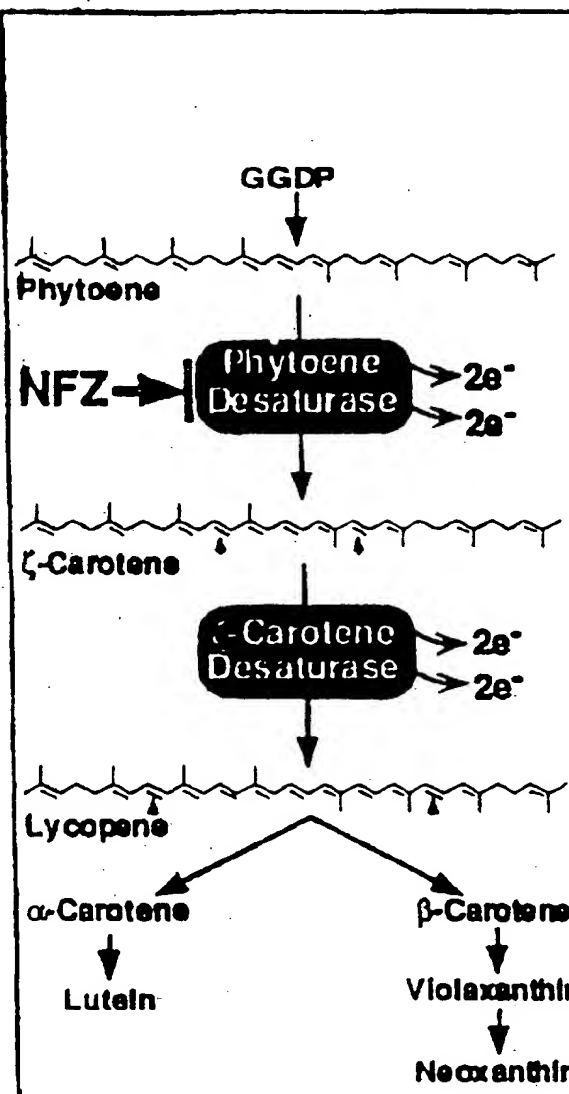
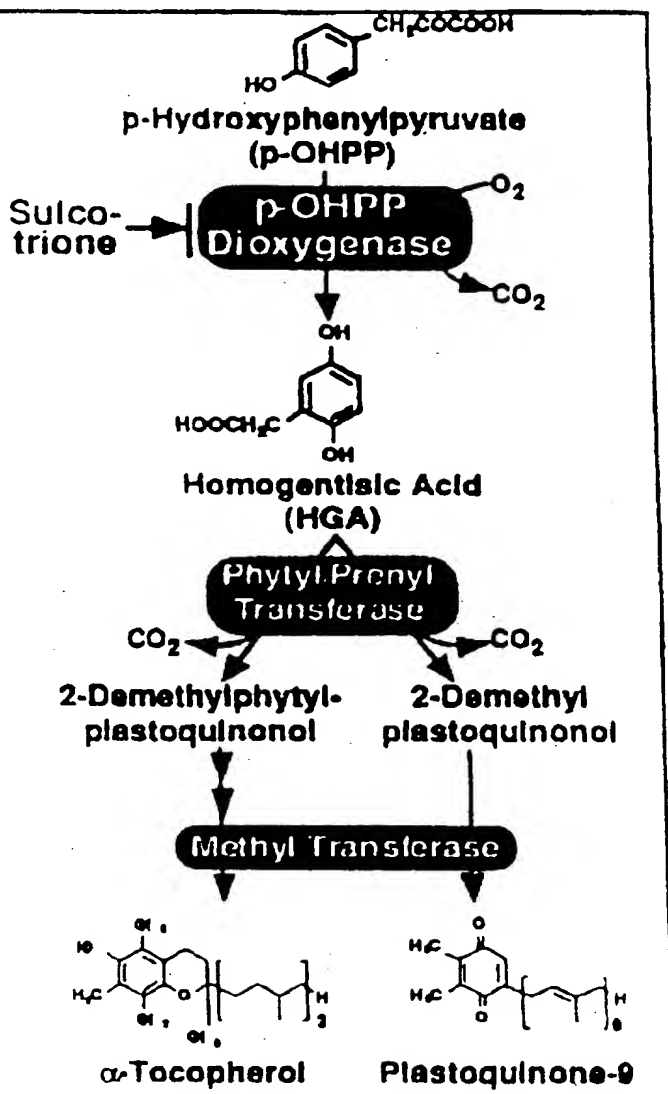
Carotenoid PathwayTocopherol/Plastoquinone Pathway

FIG 1

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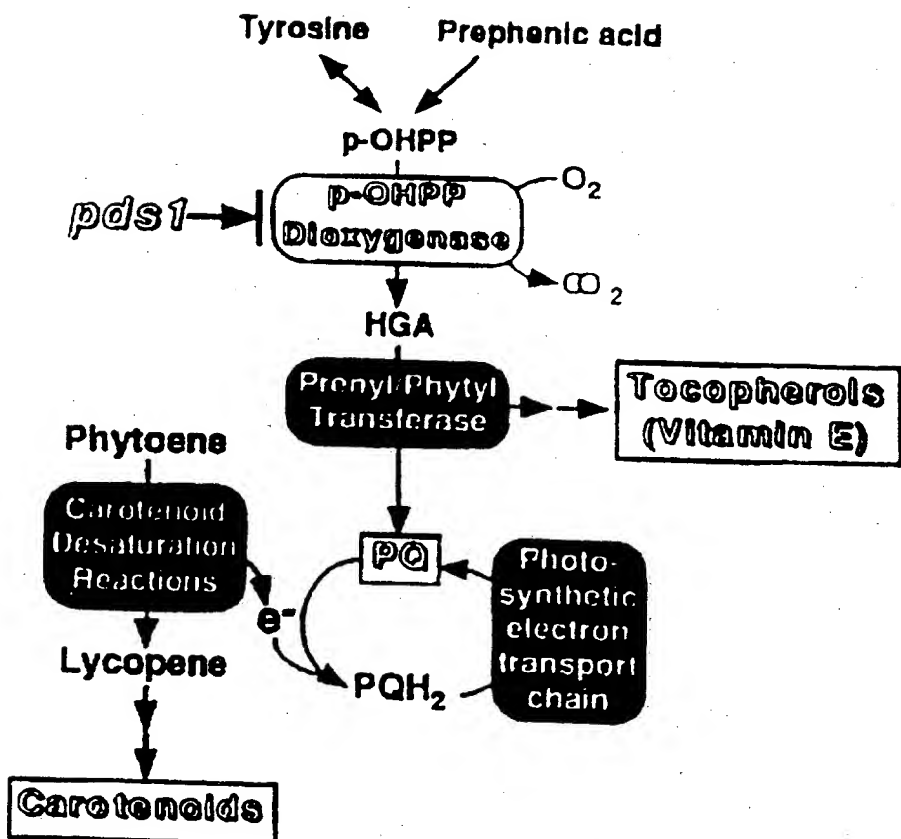


FIG 2

3/6

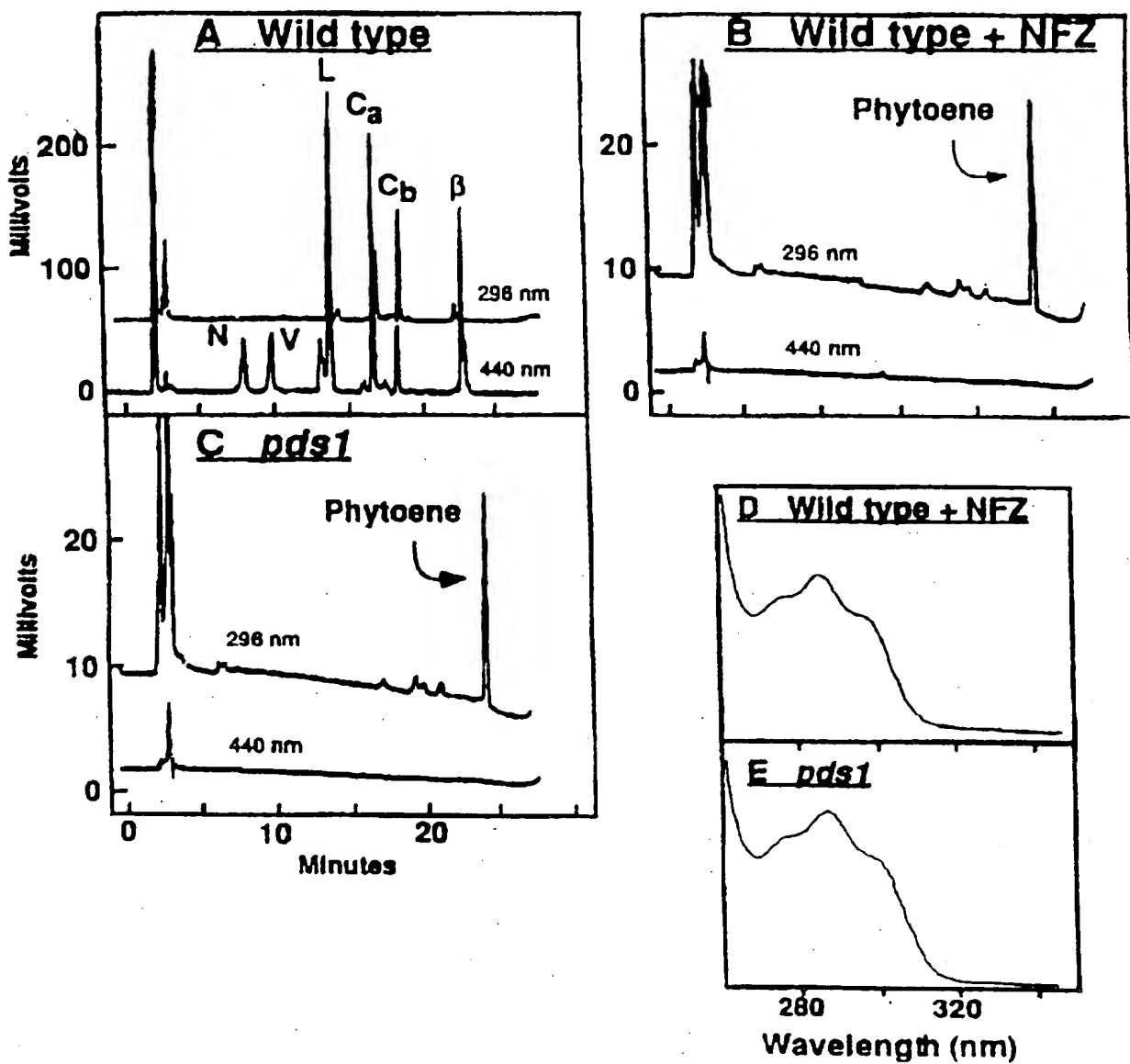


FIG 3A - 3E

4/6

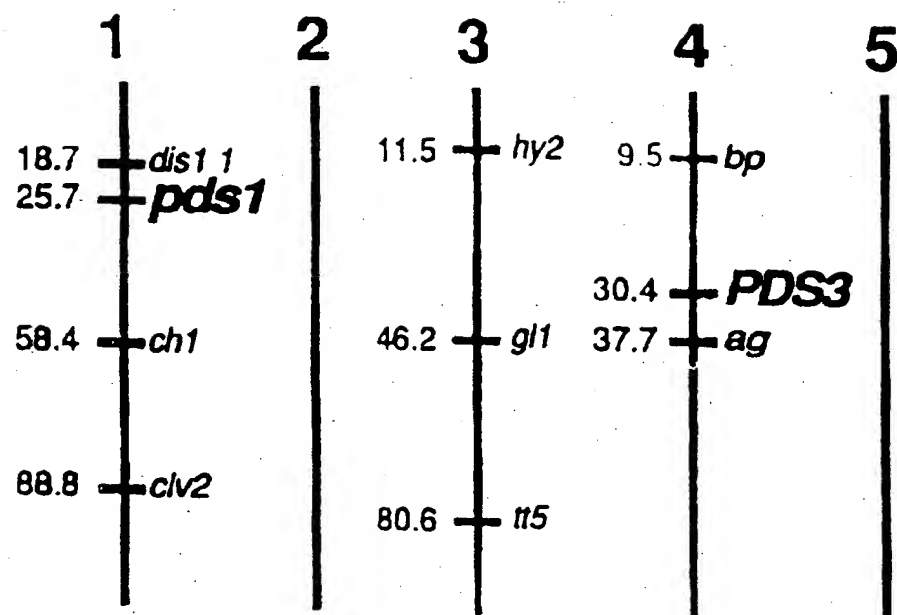


FIG 4

5/6

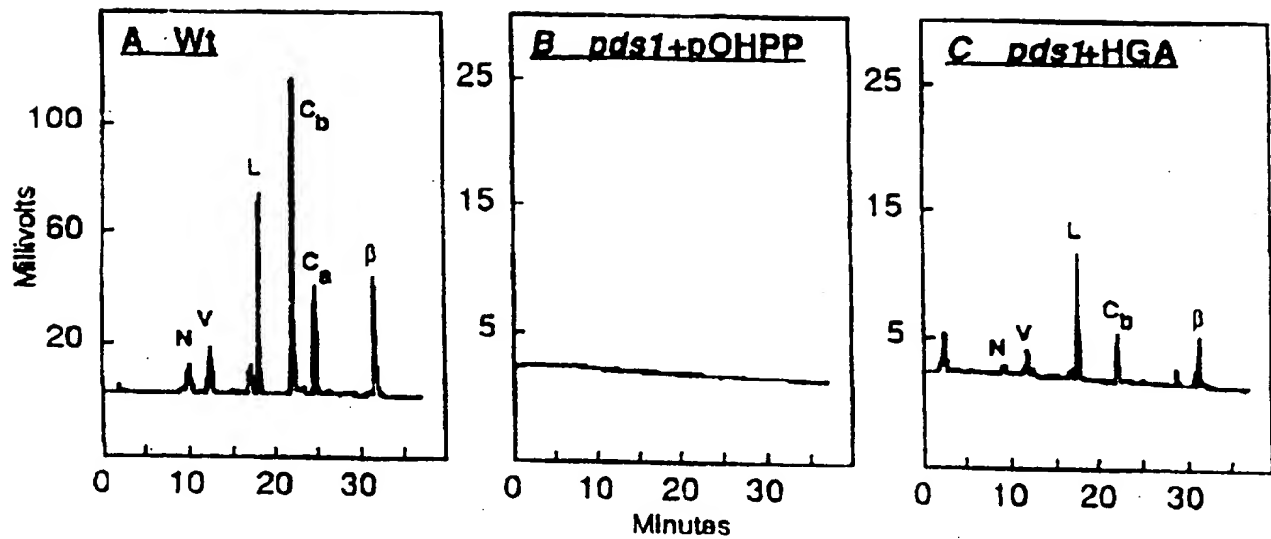
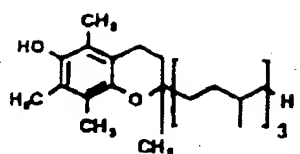
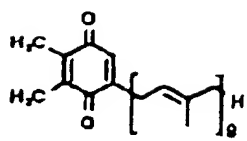


FIG 5A - 5C

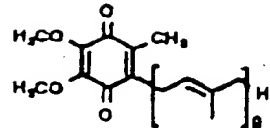
6/6



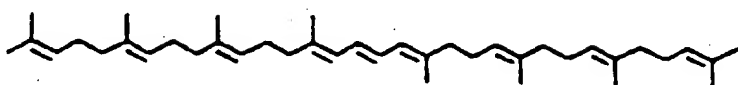
α -Tocopherol (1a)



Plastoquinone-9 (4)



Ubiquinone-9 (3)



Phytoene (2)

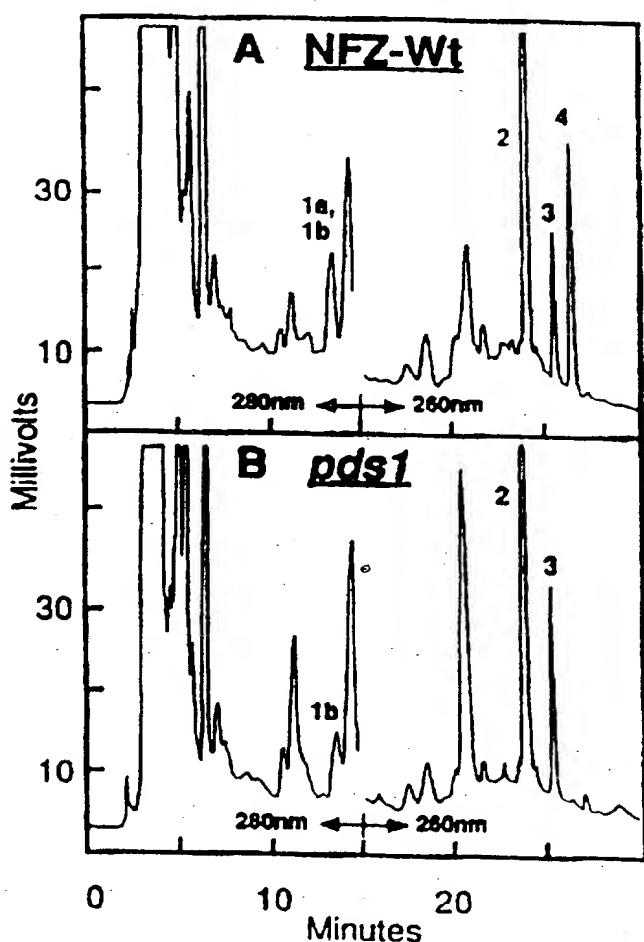


FIG 6A - 6B

INTERNATIONAL SEARCH REPORT

International application No.
US97/01384

A. CLASSIFICATION OF SUBJECT MATTER

IPC(6) : C12N 1/20, 15/00, 15/63; C07H 21/04; A01H 1/00, 1/02
US CL : 435/172.1, 252.3, 320.1, 419, 423; 536/23.2, 23.6; 800/205, 250

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 435/172.1, 252.3, 320.1, 419, 423; 536/23.2, 23.6; 800/205, 250

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

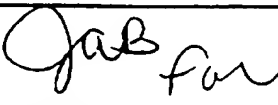
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

Please See Extra Sheet.

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	ENDO et al. Primary Structure Deduced from Complementary DNA Sequence and Expression in Cultured Cells of Mammalian 4-Hydroxyphenylpyruvic Acid Dioxygenase. The Journal of Biological Chemistry. 05 December 1992. Vol. 267, No. 34, pages 24235-24240, especially page 24238.	1-17
Y	NEWMAN et al. Genes Galore: A Summary of Methods for Accessing Results from Large-Scale Partial Sequencing of Anonymous Arabidopsis cDNA Clones. Plant Physiology. December 1994. Vol. 106, No. 4, pages 1241-1255, see entire document.	1-17

<input type="checkbox"/>	Further documents are listed in the continuation of Box C.	<input type="checkbox"/>	See patent family annex.
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O document referring to an oral disclosure, use, exhibition or other means			
P document published prior to the international filing date but later than the priority date claimed		*&*	document member of the same patent family

Date of the actual completion of the international search	Date of mailing of the international search report
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INTERNATIONAL SEARCH REPORT

International application No.

PCT/US97/01384

B. FIELDS SEARCHED

Electronic data bases consulted (Name of data base and where practicable terms used):

APS, BIOSIS, AGRICOLA, MPSEARCH (sequences only)

search terms: hydroxyphenyl pyruvic acid, DNA, clon?, SEQ ID NO:1 and NO:2

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(21) International Application Number: PCT/US97/01384 (22) International Filing Date: 28 January 1997 (28.01.97)		(81) Designated States: AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, UZ, VN, ARIVO patent (KE, LS, MW, SD, SZ, UG), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG).	
(30) Priority Data: 08/592,900 29 January 1996 (29.01.96) US		Published <i>With international search report.</i>	
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(54) Title: CLONED PLANT P-HYDROXYPHENYL PYRUVIC ACID DIOXYGENASE			
(57) Abstract <p>A cDNA clone from <i>Arabidopsis thaliana</i>, pHPP1.5, SEQ ID NO:1, which encodes the enzyme p-hydroxyphenyl pyruvic acid dioxygenase, is disclosed. A vector and microbial host containing a DNA sequence coding for the expression of <i>Arabidopsis thaliana</i> p-hydroxyphenyl pyruvic acid dioxygenase, and a genetic construct containing a DNA sequence coding for the expression of <i>Arabidopsis thaliana</i> p-hydroxyphenyl pyruvic acid dioxygenase, together with a promoter located 5' to the DNA coding sequence and a 3' termination sequence, are also disclosed. A method of creating a transgenic plant in which production of plastoquinones, vitamin E, and carotenoids has been modified, is also disclosed.</p>			

* (Referred to in PCT Gazette No. 43/1997, Section II)

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5 CLONED PLANT P-HYDROXYPHENYL PYRUVIC ACID DIOXYGENASE

This invention was made with Government support under Grant Number 93373069083 awarded by the U.S. Department of Agriculture. The Government has certain rights in the invention.

10

Field Of The Invention

The present invention relates to a molecular approach for modifying the synthesis of vitamin E, plastoquinone, and carotenoids in plants by use of a full-length cloned cDNA which encodes a p-hydroxyphenyl pyruvic acid dioxygenase enzyme.

Background Of The Invention

The chloroplasts of higher plants contain many unique, interconnected biochemical pathways that produce an array of secondary metabolite compounds which not only perform vital functions within the plant but are also important from agricultural and nutritional perspectives. Three such secondary metabolites are the lipid soluble, chloroplastically synthesized compounds vitamin E (α -tocopherol or α -toc), plastoquinones (PQ), and carotenoids, which together perform many crucial biochemical functions in the chloroplast. PQ and vitamin E are quinone compounds synthesized by a common pathway in the plastid; carotenoids are tetraterpenoids synthesized by a separate plastid-localized pathway.

Plastoquinone (PQ) often accounts for up to 50% of the total plastidic quinone pool in green tissues. The primary function of PQ is as a fundamental component of the photosynthetic electron transport chain, acting as an electron carrier between

photosystem II and the cytochrome b_6f complex. PQ likely has other less well studied functions in plastids, namely in acting as a direct or intermediate electron carrier for a variety of other biosynthetic reactions in the chloroplast.

Vitamin E is the second major class of chloroplastic quinones, accounting for up to 40% of the quinone pool in plastids. The essential nutritional value of tocopherols was recognized around 1925, and the compound responsible for Vitamin E activity was first identified as α -tocopherol in 1936. α -Toc has a well-documented role in mammals as an antioxidant, and a similar, though less well understood antioxidant role in plants. Liebler, et al., Toxicology 23:147-169, 1993; Hess, Anti-oxidants in Higher Plants, CRC Press: 111-134, 1993.

Carotenoids are a separate, diverse group of lipophilic pigments synthesized in plants, fungi, and bacteria. In photosynthetic tissues, carotenoids function as accessory pigments in light harvesting and play important roles in photo-protection by quenching free radicals, singlet oxygen, and other reactive species. Siefermann-Harms, Physiol. Plantarum. 69:561-568, 1987. In the plastids of non-photosynthetic tissues, high levels of carotenoids often accumulate providing the intense orange, yellow, and red coloration of many fruits, vegetables, and flowers (Pfander, Methods in Enzym., 213A, 3-13, 1992). In addition to their many functions in plants, carotenoids and their metabolites also have important functions in animals, where they serve as the major source of Vitamin A (retinol), and have been identified as providing protection from some forms of cancer due to their antioxidant activities. Vitamin E's antioxidant activities are also thought to protect against some forms of cancer, and may act synergistically with carotenoids in this regard.

Liebler, et al., Toxicology 23:147-169, 1993; Krinsky, J. Nutr. 119:123-126, 1989.

Tocopherol and Plastoquinone Synthesis

α -Tocopherol and plastoquinone are the most abundant quinones in the plastid and are synthesized by the common pathway shown in Figure 1. The precursor molecule for both compounds, homogentisic acid (HGA), is produced in the chloroplast from the shikimic acid pathway intermediate p-hydroxyphenyl pyruvic acid (pOHPP), in an oxidation/decarboxylation reaction catalyzed by the enzyme p-hydroxyphenyl pyruvic acid dioxygenase (pOHPP dioxygenase). Homogentisic acid is subject to phytylation/prenylation (phytylpyrophosphate and solanylpyrophosphate, C₂₀ and C₄₅, respectively) coupled to a simultaneous decarboxylation by a phytyl/prenyl transferase to form the first true tocopherol and plastoquinone intermediates, 2-demethylphytylplastoquinol and 2-demethylplastoquinol-9, respectively. A single ring methylation occurs on 2-demethylplastoquinol to yield plastoquinol-9 (PQH₂) which is then oxidized to plastoquinone-9 (PQ). This oxidation is reversible and is the basis of electron transport by plastoquinone in the chloroplast.

The preferred route, as established in spinach, for α -tocopherol formation from 2-demethylphytylplastoquinol appears to be 1) ring methylation of the intermediate, 2- α -demethylphytylplastoquinol, to yield phytylplastoquinol, 2) cyclization to yield d-tocopherol and, finally, 3) a second ring methylation to yield α -tocopherol. Ring methylation in both tocopherol and plastoquinone synthesis is carried out by a single enzyme that is specific for the site of methylation on the ring, but has

relatively broad substrate specificity and accommodates both classes of quinone compounds. This methylation enzyme is the only enzyme of the pathway that has been purified from plants to date.

5 d'Harlingue, et al., J.Biol.Chem. 26:15200, 1985. All enzymatic activities of the α -toc/PQ pathway have been localized to the inner chloroplast envelope by cell fractionation studies except for pOHPP dioxygenase and the tocopherol cyclase enzyme. Difficulties with cell
10 fractionation methods, low activities for some of the enzymes, substrate stability and availability and assay problems, make studying the pathway biochemically difficult.

Vitamin E and PQ levels, ratios, and total amounts vary by orders of magnitude in different plants, tissues and developmental stages. Such variations indicate that the vitamin E and PQ pathway is both highly regulated and has the potential for manipulation to modify the absolute levels and ratios 20 of the two end products. The pathway in Figure 1 makes it clear that production of homogentisic acid by pOHPP dioxygenase is likely to be a key regulatory point for bulk flow through the pathway, both because HGA production is the first committed step in α -toc/PQ synthesis, and also because the reaction is essentially irreversible. Therefore modifying the levels of HGA by modifying pOHPP dioxygenase activity should have a direct impact on the total α -toc/PQ biosynthetic accumulation in plant tissues, and, as 25 described below, because of the connection of PQ and carotenoid synthesis, should also affect carotenoid synthesis in plant tissues.

Carotenoid Biosynthesis; Quinones as Electron Carriers

In plants, carotenoids are synthesized and 35 accumulate exclusively in plastids via the pathway shown on the left-hand side of Figure 1. The first

committed step in carotenoid synthesis is the condensation of two molecules of the C₂₀ hydrocarbon geranylgeranyl pyrophosphate (GGDP) by the enzyme phytoene synthase, to form the colorless C₄₀ hydrocarbon, phytoene. In oxygenic photosynthetic organisms (e.g. plants, algae, and cyanobacteria), phytoene undergoes two sequential desaturation reactions, catalyzed by phytoene desaturase, to produce ζ -carotene through the intermediate phytofluene. Subsequently, ζ -carotene undergoes two further desaturations, catalyzed by ζ -carotene desaturase, to yield the red pigment lycopene. Lycopene is cyclized to produce either α -carotene or β -carotene, both of which are subject to various hydroxylation and epoxidation reactions to yield the carotenoids and xanthophylls most abundant in photosynthetic tissues of plants, lutein, β -carotene, violaxanthin and neoxanthin.

The genes encoding the first two enzymes of the carotenoid pathway (phytoene synthase and phytoene desaturase) have been isolated and studied from a number of plant and bacterial sources in recent years. Sandmann, Eur. J. Biochem. 223:7-24, 1994. Phytoene desaturase has been the most intensively studied, both because it is a target for numerous commercially important herbicides, and also because the phytoene desaturation reaction is thought to be a rate limiting step in carotenoid synthesis. Molecular and biochemical studies suggest that two types of phytoene desaturase enzymes have evolved by independent evolution: the crtI-type found in anoxygenic photosynthetic organisms (e.g. *Rhodobacter* and *Erwinia*), and the pds-type found in oxygenic photosynthetic organisms. Despite their differences in primary amino acid sequence, all phytoene desaturase enzymes contain a dinucleotide binding domain (FAD or NAD/NADP), which in *Capsicum annuum* has

been shown to be FAD. Hugueney et al., Eur. J. Biochem. 209:399-407, 1992. Presumably, the bound dinucleotide in both types of phytoene desaturase enzymes is reduced during desaturation and reoxidized by an unknown reductant present in the plastid or bacterium.

Several lines of evidence have suggested a role for quinones in the phytoene desaturation reaction in higher plants. Using isolated daffodil chromoplasts, Mayer and co-workers demonstrated that in an anaerobic environment, oxidized artificial quinones were required for the desaturation of phytoene while reduced quinones were ineffective. Mayer et al., Eur. J. Biochem. 191:359-363, 1990. Further supporting evidence comes from studies with the triketone class of herbicides (e.g. Sulcotrione), which cause phytoene accumulation in treated tissues but unlike the well-studied pyridazone class (e.g. Norflorazon (NFZ)) do not directly affect the phytoene desaturase enzyme. Rather, triketone herbicides competitively inhibit pOHPP dioxygenase, an enzyme common to the synthesis of both plastoquinone and tocopherols, suggesting that one or more classes of quinones may play a role in carotenoid desaturation reactions. Schulz et al., FEBS 318:162-166, 1993; Secor, Plant Physiol. 106: 1429-1433; Beyer et al., IUPAC Pure and Applied Chemistry 66:1047-1056, 1994.

Despite the well-studied, wide-spread importance of vitamin E, plastoquinone, and carotenoids to human nutrition, agriculture, and biochemical processes within plant cells, much remains unclear about their biosynthesis and accumulation in plant tissues. This uncertainty has in turn limited the potential for manipulation of the synthesis and levels of these important compounds in plants.

Summary of the Invention

In one embodiment, this invention provides a biologically pure sample of DNA which DNA comprises a sequence coding for the expression of *Arabidopsis thaliana* p-hydroxyphenyl pyruvic acid dioxygenase.

5 In other embodiments, this invention provides a vector and microbial host containing a DNA sequence sufficiently homologous to SEQ ID NO:1 so as to code for the expression of *Arabidopsis thaliana* p-hydroxyphenyl pyruvic acid dioxygenase, and a genetic construct containing a DNA sequence sufficiently homologous to SEQ ID NO:1 so as to code for the expression of *Arabidopsis thaliana* p-hydroxyphenyl pyruvic acid dioxygenase, together with a promoter located 5' to the DNA coding sequence and a 3' termination sequence.

10 15 In another embodiment, this invention provides a method of creating a transgenic plant in which the levels of the pOHPP dioxygenase enzyme are elevated sufficient such that production of plastoquinones, vitamin E, and carotenoids are modified.

20 25 It is an object of the present invention to genetically engineer higher plants to modify the production of plastoquinones, vitamin E, and carotenoids.

30 It is another object of the invention to provide transgenic plants that would express elevated levels of the pOHPP dioxygenase enzyme which would have resultant elevated resistance to the triketone class of herbicides (i.e. sulcotriione).

35 It is another object of the present invention to provide a method for the preparation of the enzyme p-hydroxyphenyl pyruvic acid dioxygenase (pOHPP dioxygenase), an enzyme which can be used to identify new pOHPPdioxygenase-inhibiting herbicides.

Other features and advantages of the invention will be apparent from the following description of the

preferred embodiments thereof and from the claims.

Brief Description of the Drawings

Fig. 1 is a diagram of the pathways for synthesis
of carotenoids, vitamin E (tocopherol), and
5 plastoquinone.

Fig. 2 is a diagram of the interconnections of
the pathways illustrated in Fig. 1.

Fig. 3A-3E are graphs of pigment analyses of
wild-type, NFZ-wt, and *pds1* tissues.

10 Fig. 4 is a physical map of the *pds1* mutation
relative to visible markers.

15 Figs. 5A-5C present the results of C18 HPLC
separation of lipid soluble pigments from wild-type
plants on MS2 media, homozygous *pds1* mutants on MS2
media supplemented with pOHPP, and homozygous *pds1*
mutants on MS2 media supplemented with homogentistic
acid (HGA).

20 Figs. 6A-6B present the results of C8 HPLC
analyses of quinones in NFZ-wt and *pds1* tissues.

Detailed Description Of The Invention

As described above, both Vitamin E,
plastoquinones and carotenoids are synthesized and
accumulated in plastids by the pathways shown in
Figure 1. This specification describes the
identification, isolation, characterization and
functional analysis of a higher plant pOHPP
dioxygenase cDNA, its role in α -toc, PQ and carotenoid
synthesis, and the use of this cDNA to modify pOHPP
dioxygenase activity in plant tissues and hence the
30 accumulation of one or more of the compounds
plastoquinones, vitamin E, and carotenoids in plant
tissues. The overexpression of pOHPP dioxygenase in
transgenic plants will modify the enzyme-to-inhibitor
ratio of plant tissues exposed to triketone
35 herbicides, as compared to non-transgenic plants,

resulting in increased herbicide resistance. The present specification also describes a genetic construct for use in the production of pOHPP dioxygenase, an enzyme useful in identifying new pOHPP dioxygenase-inhibiting herbicides.

By genetic analysis the present inventors have shown that the vitamin E, plastoquinone, and carotenoid biosynthetic pathways are interconnected and share common elements as shown in Figure 2. From mutational studies in *Arabidopsis thaliana*, the present inventors identified one genetic locus, designated *pds1* (*pds*= phytoene desaturation), the disruption of which results in accumulation of the first carotenoid of the carotenoid biosynthetic pathway, phytoene. Surprisingly, though this mutation disrupts carotenoid synthesis and was originally identified on this basis, it does not map to the locus encoding the phytoene desaturase enzyme. Evidence indicates that *pds1* defines a second gene product in addition to the phytoene desaturase enzyme, necessary for phytoene desaturation and hence carotenoid synthesis in higher plants. This gene product proved to be pOHPP dioxygenase.

To provide a molecular mechanism for manipulating synthesis and accumulation of the compounds plastoquinone, vitamin E, and carotenoids, the present inventors used a molecular genetic approach, taking advantage of the model plant system *Arabidopsis thaliana* to define, isolate and study genes required for synthesis of the compounds in plants. The flowering plant *Arabidopsis thaliana* has come into wide use as a model system to explore the molecular biology and genetics of plants. *Arabidopsis* offers many advantages for genetic analysis: it can be selfed and very large numbers of progeny can be obtained (up to 10,000 seeds from a single plant). Furthermore, *Arabidopsis* has a short generation time of five to six

weeks, so crosses can be set up and the progeny analyzed within reasonable periods of time. Mutation screens have identified thousands of mutations affecting many aspects of basic plant biology, 5 including morphogenesis, photosynthesis, fertility, starch and lipid metabolism, mineral nutrition, and so on. In addition, its haploid genome is only about 10^8 base pairs.

An important aspect of the successful approach 10 used here is that essential components were first functionally defined genetically, prior to their isolation, analysis and molecular manipulation. Briefly, potential mutants were identified by a combination of phenotypic and biochemical screening, 15 characterized at the genetic and molecular levels, loci of interest selected, and the corresponding genes then cloned and studied further. By this approach, the inventors genetically defined and isolated cDNAs for one gene, *pds1*, whose mutation disrupts synthesis 20 of all three classes of compounds in the plastid, tocopherols, plastoquinones and carotenoids. Based on biochemical analysis of the *pds1* mutant, the *pds1* gene was identified as affecting the activity of pOHPP 25 dioxygenase, a crucial enzyme of the plastidic quinone pathway in plants (Figure 1), that is directly required for the synthesis of plastoquinone and α -tocopherol and indirectly for carotenoid synthesis. In particular, the deduced function of the *pds1* mutant and pOHPP dioxygenase enzyme are noted in Figure 2.

The present inventors demonstrated by biochemical complementation that the *pds1* mutation affects the enzyme p-hydroxyphenyl pyruvic acid dioxygenase (pOHPP dioxygenase), because *pds1* plants can be rescued by growth on the product but not the substrate of this 30 enzyme, homogentisic acid (HGA) and p-hydroxyphenylpyruvate (pOHPP), respectively. pOHPP dioxygenase is the key branch point enzyme and

committed step in the synthesis of both Vitamin E and plastoquinones and several independent lines of biochemical evidence confirm *pds1* affects this enzyme (Figures 1, 5, 6). These results provide the first 5 genetic evidence that plastoquinones are essential components for carotenoid synthesis in higher plants, most likely as an electron carrier/redox element in the desaturation reaction (Figure 2). The *Arabidopsis* pOHPP dioxygenase gene/cDNA thus provides a basis for 10 modifying the production of plastoquinones, α -tocopherol and carotenoids in all higher plants.

Specifically, the specification describes the 15 genetic identification of the *Arabidopsis* pOHPP dioxygenase gene by mutational analysis, the physical isolation and functional confirmation of an *Arabidopsis* pOHPP dioxygenase cDNA, its nucleotide sequence and its use to isolate pOHPP dioxygenase genes and cDNAs from other plant species. Also included in the specification is a description of the 20 use of the *Arabidopsis* pOHPP dioxygenase cDNA, and related cDNAs from other plants, to positively or negatively modify the expression/activity of pOHPP dioxygenase by recombinant techniques (overexpression, cosuppression, antisense, etc.) in any and all plant 25 tissues, especially leaf and fruit tissues, to positively or negatively affect the production of α -toc, PQ and carotenoids.

Elevating pOHPP dioxygenase protein levels 30 increases the amount of homogentisic acid (HGA) synthesized in plant tissues. Because HGA is the limiting precursor molecule for α -toc and PQ synthesis (the end products of the pathway), increasing HGA synthesis increases the levels of α -toc (Vitamin E) and PQ in plant tissues. The increase in PQ 35 indirectly increases the synthesis of carotenoids, which require PQ for their synthesis. In addition, the increase in PQ increases photosynthetic efficiency

by increasing electron flow between photosystem II and photosystem I, because PQ is the primary electron transporter between the two photosystems. The increase in α -toc, a well-studied antioxidant in mammals, increases the ability of plants to withstand oxidative stresses, such as that caused by high light, high temperature, water stress, ozone stress, UV stress or other abiotic or biotic stresses. Elevating the levels of pOHPP dioxygenase will modify the dose response curve of herbicides targeting pOHPP dioxygenase, thus increasing the relative resistance to such herbicides in transgenic plants as compared to native plants of the same species. Inhibiting the expression of pOHPP dioxygenase is expected to have the opposite effect.

Genetic Construct

To express pOHPP dioxygenase in a plant, it is required that a DNA sequence containing the pOHPP dioxygenase coding sequence be combined with regulatory sequences capable of expressing the coding sequence in a plant. A number of effective plant promoters, both constitutive and developmentally or tissue specific, are known to those of skill in the art. A transcriptional termination sequence (polyadenylation sequence) may also be added. Plant expression vectors, or plasmids constructed for expression of inserted coding sequences in plants, are widely used in the art to assemble chimeric plant expression constructs including the coding sequence, and to conveniently transfer the constructs into plants. A sequence which codes for pOHPP dioxygenase includes, for example, SEQ ID NO:1, or versions of the designated sequence sufficient to effect coding for the expression of pOHPP dioxygenase. Commonly used methods of molecular biology well-known to those of skill in the art may be used to manipulate the DNA

sequences.

By "genetic construct" we mean any of a variety of ways of combining the protein-encoding sequences with a promoter sequence (and termination sequence, if necessary) in a manner that operably connects the promoter sequence (and termination sequence, if present) with the protein-encoding sequences.

Typically, the promoter sequence will be "upstream" of a protein-encoding sequence, while the termination sequence, if used, will be "downstream" of the protein-encoding sequences.

The protein-encoding, promoter and termination sequences may be combined on a plasmid or viral vector, and inserted into a microbial host. Other functional sequences may be added to the gene construct. Alternatively, the protein-encoding, promoter, and termination sequence, if added, may be combined with any other needed functional sequences and used without a vector.

The DNA sequence described by SEQ ID NO:1 is sufficient to effect coding for the expression of pOHPP dioxygenase. However, it is envisioned that the above sequence could be truncated and still confer the same properties. It is not known at present which specific deletions would be successful, but it is likely that some deletions to the protein would still result in effective enzymatic activity. One skilled in the art of molecular biology would be able to take the designated sequence and perform deletional analysis experiments to determine what portions of the designated sequence are essential to effect coding for the expression of pOHPP dioxygenase. One could create a genetic construct with the candidate deletion mutations and perform experiments as described below in the Examples, to test whether such deletion mutation sequences effect coding for the enzyme. Expression of the enzyme activity indicates a

successful deletion mutant or mutants. In this manner, one could determine which parts of the designated sequence is essential for expression of the enzyme.

5 It is also known that the genetic code is degenerate, meaning that more than one codon, or set of three nucleotides, codes for each amino acid. Thus it is possible to alter the DNA coding sequence to a protein, such as the sequence for pOHPP dioxygenase
10 described here, without altering the sequence of the protein produced. Selection of codon usage may affect expression level in a particular host. Such changes in codon usage are also contemplated here.

15 It is further contemplated that using the *Arabidopsis* pOHPP gene coding sequence described here, that the homologous pOHPP dioxygenase sequences from other higher plants can be readily recovered. Oligonucleotides can be made from the sequence set forth below to either hybridize against cDNA or
20 genomic libraries or used for PCR amplification of homologous pOHPP dioxygenase sequences from other plants.

Once a pOHPP gene is in hand, whether from *Arabidopsis* or from some other plant species, it then becomes possible to insert a chimeric plant expression 25 genetic construct into any plant species of interest. Suitable plant transformation methods exist to insert such genetic constructs into most, if not all, commercially important plant species. Presently known
30 methods include *Agrobacterium*-mediated transformation, coated-particle gene delivery (Biolistics) and electroporation, in which an electric voltage is used to facilitate gene insertion. All these methods, and others, can insert the genetic construct into the genome of the resulting transgenic plant in such a way
35 that the genetic construct becomes an inheritable trait, transmitted to progeny of the original

transgenic plant by the normal rules of Mendelian inheritance. Thus, once a genetic construct expressing a pOHPP gene is inserted into a plant, it can become a part of a plant breeding program for transfer into any desired genetic background.

To over-express pOHPP dioxygenase, a genetic construct may be used with a higher strength promoter. To inhibit expression of endogenous pOHPP dioxygenase, an antisense genetic construct can be made, as is known by those of skill in the art, to reduce the level of pOHPP dioxygenase present in the plant tissues.

EXAMPLES

Isolation of *pds1*, a mutant defective in carotenoid synthesis

To further understand carotenoid biosynthesis and its integration with other pathways in the chloroplast in higher plants, the present inventors studied the pathway by isolating *Arabidopsis thaliana* mutants that are blocked in carotenoid synthesis.

Plants homozygous for defects in the early stages of carotenoid synthesis (e.g. prior to production of β -carotene) are lethal when grown in soil and the isolation of such mutations requires the design of screening procedures to identify plants heterozygous for soil lethal mutations. The present inventors found that most soil lethal, homozygous pigment-deficient *Arabidopsis* mutants can be grown to near maturity in tissue culture on Murashige and Skoog basal media (Murashige and Skoog, *Physiol. Plant.* 15:473-497, 1962) supplemented with sucrose (MS2 media). Under these conditions, photosynthesis and chloroplast development are essentially dispensable and all the energy and nutritional needs of the plant are supplied by the media.

Greater than 500 lines from the 10,000 member

Feldmann T-DNA tagged *Arabidopsis thaliana* population (Forsthoefel et al., Aust. J. Plant Physiol. 19:353-366, 1992) were selected for pigment analysis based on their segregation for lethal pigment mutations. Seed from plants heterozygous for lethal pigment mutations were surface sterilized, grown on MS2 media, the segregating pigment mutants identified, tissue harvested from individual plants, and HPLC pigment analysis performed. Although numerous mutant lines with severe pigment deficiencies were identified, only two were found to be carotenoid biosynthetic mutants. One mutant line isolated from this group, *pds1*, is described in detail here.

The hallmark phenotype for disruption of a biosynthetic pathway is the accumulation of an intermediate compound prior to the site of blockage. Such blockage of the carotenoid pathway can be mimicked chemically by treatment of wild-type plants with the herbicide NFZ, an inhibitor of the phytoene desaturase enzyme (Figure 1) which has been reported to cause accumulation of phytoene in treated tissues. Britton, Z. Naturforsch 34c:979-985, 1979. Figs. 3A-3E present the results of pigment analysis of wild-type, NFZ-wt, and *pds1* tissues. Abbreviations in Figs. 3A-3E are as follows: N, neoxanthin; V, violaxanthin; L, lutein; Cb, chlorophyll b; Ca, chlorophyll a; β, β-carotene.

Figure 3A shows C_{18} Reverse Phase HPLC analysis of the carotenoids that accumulate in wild-type *Arabidopsis thaliana* leaves. In comparison, Figure 3B shows the pigment profile for NFZ treated wild-type (NFZ-Wt). Spectral analysis of the strongly absorbing 296nm peak at 33 minutes in NFZ-Wt tissue shows absorbance maxima at 276, 286, and 298nm, indicative of phytoene (Figure 3D). Figure 3C shows pigment analysis of tissue culture grown homozygous *pds1* mutant plants. The low absorbance at 440nm in Figures

3B and C demonstrates that like NFZ-Wt, *pds1* mutants lack all chlorophylls and carotenoids that normally accumulate in wild-type tissue (compare to Figure 3A). However, unlike wild-type, *pds1* mutants contain a peak with a retention time at approximately 33 minutes that absorbs strongly at 296nm. The retention time and absorbance of the 33-minute peak in the *pds1* mutant corresponds to the phytoene peak in pigment extracts of NFZ-Wt tissue (Figure 3B). Spectral analysis of the 33-minute peak from *pds1* is shown in Figure 3E and is virtually identical to the spectra of phytoene from NFZ-Wt tissue (Figure 3D) as well as to the published spectra for phytoene. These results confirm the chemical identity of the accumulating compound in *pds1* as phytoene and conclusively demonstrate that the *pds1* mutation disrupts carotenoid biosynthesis.

Carotenoid Analysis

For quantitative and qualitative carotenoid analysis, plant tissue is placed in a microfuge tube and ground with a micropesle in 200 μ l of 80% acetone. 120 μ l of ethyl acetate is added and the mixture vortexed. 140 μ l of water is added and the mixture centrifuged for 5 minutes. The carotenoid containing upper phase is then transferred to a fresh tube and vacuum dried in a Jouan RC1010 Centrifugal Evaporator. The dried extract is resuspended in ethyl acetate at a concentration of 0.5mg fresh weight of tissue per μ l and either analyzed immediately by HPLC or stored at -80°C under nitrogen.

Carotenoids were separated by reverse-phase HPLC analysis on a Spherisorb ODS2 5 micron C₁₈ column, 25 cm in length (Phase Separations Limited, Norwalk, CT) using a 45 minute gradient of Ethyl Acetate (0-100%) in Acetonitrile/water/triethylamine (9:1:0.01 v/v), at a flow rate of 1 ml per minute (Goodwin and Britton, 1988). Carotenoids were identified by retention time

relative to known standards with detection at both 296nm and 440nm. When needed, absorption spectra for individual peaks were obtained with a Hewlett Packard 1040A photodiode array detector and compared with 5 published spectra or available standards.

Quinone analysis

Quinones were extracted from tissue using a method modified from that described in Bligh et al., Can. J. Biochem. Physiol. 37:911-917, 1959. Frozen 10 plant tissue was ground in a mortar with 3 volumes of chloroform and 6 volumes of methanol and transferred to a test tube. Water and additional chloroform were added until a biphasic mixture was obtained. The quinone containing chloroform phase was then 15 collected. To increase yields, the aqueous phase was back-extracted with chloroform, the two chloroform phases pooled, and then filtered through Whatman #3 filter paper. The resulting filtrate was dried under a constant stream of nitrogen. Once dried, the pellet 20 was resuspended in methanol at a concentration of 10mg fresh weight per ml and immediately analyzed by HPLC. Quinones were resolved by reversed-phase HPLC analysis on a LiChrosorb RP-8, 5 micron column, 25cm in length, (Alltech, San Jose, CA) using an isocratic solvent of 25 10% H₂O in Methanol for the first 14 minutes, at which time the solvent was switched to 100% methanol for the remainder of the run (modified from the method described in Lichtenthaler, Handbook of Chromatography, CRC Press, 115-159, 1984). The flow 30 rate was 1ml per minute for the duration. Peaks were identified based upon the retention time of known standards with detection at 280nm for α -tocopherol and 35 260nm for plastoquinone and ubiquinone as well as by absorption spectra from a Hewlett Packard 1040A photodiode array detector. When needed, fractions represented by individual chromatographic peaks were

collected, and submitted to the Southwest Environmental Health Science Center, Analytical Core laboratory for mass spectral analysis. Results were obtained using a TSQ7000 tandem mass spectrometer (Finnigan Corp., San Jose, CA) equipped with an atmospheric pressure chemical ionization source operated in the positive ion mode. The instrument was set to unit resolution and the samples were introduced into the source in a 0.3 ml/minute methanol stream and ionized using a 5kV discharge.

Genetic analysis of *pds1*

The genetic nature of the *pds1* mutation was determined by analyzing seeds resulting from selfing *pds1* heterozygous plants. Prior to desiccation, F1 seeds were scored as either green (wild-type or heterozygous) or white (homozygous). A 3:1 segregation ratio was observed (146 green seeds: 48 white seeds), indicating that *pds1* is inherited as single recessive nuclear mutations ($\chi^2=0.01$, $p > 0.90$). Because *pds1* mutants are inhibited in the desaturation of phytoene, the inventors believed that it might be a mutation in the phytoene desaturase enzyme, which had previously been mapped to chromosome 4, between ag and bp. Wetzel et al., Plant J. 6:161-175, 1994. To test this hypothesis, the *pds1* mutation was mapped relative to visible markers. The *pds1* mutation was found to map to chromosome 1, approximately 7 cM from *dis1* toward *clv2*. Franzmann et al., Plant J. 7:341-350, 1995. These data points are summarized in Figure 4 and establish that *pds1* does not map to the phytoene desaturase enzyme locus, thus proving that the *pds1* mutation is not in the phytoene desaturase enzyme. This data provided important insight for characterization of the *pds1* mutant.

Homozygous *pds1* mutants can be rescued by Homogentisic Acid, an intermediate in plastoquinone and tocopherol biosynthesis

As described earlier, previous research suggesting a role for quinones and pOHPP dioxygenase in phytoene desaturation lead the present inventors to investigate the quinone biosynthetic pathway in the *pds1* mutant. The early stages of plastoquinone/tocopherol synthesis were functionally analyzed by growth in the presence of two intermediate compounds in the pathway, p-hydroxyphenylpyruvate (pOHPP) and homogentisic acid (HGA) (refer to Figures 1 and 2). Albino *pds1* homozygous plants were first germinated on MS2 media and then transferred to MS2 media supplemented with 100 μ M of either pOHPP or HGA. *pds1* plants remained albino when transferred to media containing pOHPP but greening occurred when *pds1* plants were transferred to media containing HGA. Figs. 5A-5C present the results of complementation of the *pds1* mutation with homogentisic acid. Each profile represents pigments extracted from 10mg fresh weight of tissue. Abbreviations used in Figs. 5A-5C are as described in Figs. 3A-3E. HPLC analysis with detection at 440nm of the carotenoids extracted from *pds1* plants grown on pOHPP and HGA are shown in Figures 5B and C, respectively. The pigment profiles of *pds1* mutants grown on pOHPP are similar to the profiles of *pds1* plants grown on MS2 media shown in Figure 3B. Comparison of the pigment profiles for *pds1* + HGA tissue and wild-type tissue (Figures 5A and 5C) indicates that growth in the presence of HGA is able to qualitatively restore a wild-type carotenoid profile to albino, homozygous *pds1* plants. These results indicate that the *pds1* mutation affects the enzyme pOHPP dioxygenase, because *pds1* mutants are not altered by growth on the substrate of this enzyme, pOHPP, but rather, are restored qualitatively to wild-type pigmentation by growth on the product of

this enzyme, HGA (refer to Figures 1 and 2). The complementation of *pds1* with HGA also indicates that intermediates or end products of this pathway (plastoquinone and/or tocopherols, refer to Figures 1 and 2) are necessary components for phytoene desaturation in plants and confirms the observation of Schultz et al. in FEBS where inhibitors of pOHPP dioxygenase were shown to cause accumulation of phytoene.

10 **HPLC analysis conclusively demonstrates that *pds1* is a mutation in the plastoquinone/ tocopherol biosynthetic pathway that also affects carotenoid synthesis**

In addition to biochemical complementation of *pds1* mutants, the plastoquinone/tocopherol pathway was also directly analyzed in *pds1* tissue by utilizing C₈ HPLC to resolve total lipid extracts and identify three separate classes of quinones: ubiquinone, plastoquinone, and α -tocopherol (Vitamin E) (Figures 5 and 6). Ubiquinone and plastoquinone perform analogous electron transport functions in the mitochondria and chloroplast, respectively, but are synthesized by different pathways in separate subcellular compartments (Goodwin et al., Introduction to Plant Biochemistry, Oxford, Pergamon Press, 1983), making ubiquinone an ideal internal control in these analyses. Figure 6 shows the C₈ HPLC analysis of lipid soluble extracts from NFZ-Wt tissue and *pds1* tissue. In NFZ-Wt tissue (Figure 6A), peaks 3 and 4 were identified as ubiquinone and plastoquinone, respectively, based on retention time (26 and 27 minutes), optical spectra, and mass spectra (results not shown). NFZ-Wt tissue contained a peak (1) with a retention time of 13.5 minutes which was identified as α -tocopherol based upon the retention time of a standard. However, optical spectroscopy and mass spectrometry demonstrated that peak 1 was composed of two major components: α -tocopherol (la) and an

unidentified compound (1b). The mass of α -tocopherol was determined to be 430 as indicated by the presence of the 431 protonated molecule while the molecular mass of the unidentified compound was 412, as indicated by the presence of the 413 protonated molecule (data not shown), clearly demonstrating the presence of two compound in peak 1. This quinone analysis demonstrates that the herbicide NFZ, which specifically inhibits the phytoene desaturase enzyme, does not affect synthesis of homogentisate derived quinones. *pds1* tissue (Figure 6B) contain ubiquinone (peak 3) but lack plastoquinone (peak 4). Additionally, though *pds1* contains a peak at 13.5 minutes, optical spectroscopy and mass spectrometry data demonstrate that this peak lacks α -tocopherol (1a) and is composed solely of the compound 1b (data not shown). Therefore, homozygous *pds1* plants accumulate ubiquinone but lack both plastoquinone and α -tocopherol. This is consistent with the *pds1* mutation affecting pOHPP dioxygenase (refer to Figures 1 and 2), as suggested by the rescue of the mutation by HGA, and provide additional evidence that the *pds1* mutation disrupts pOHPP dioxygenase.

**Isolation of a truncated, putative pOHPP dioxygenase
Arabidopsis cDNA**

The observation of Schultz et al. demonstrating that inhibitors of pOHPP dioxygenase activity disrupt carotenoid synthesis and cause accumulation of phytoene provided important insight for the characterization of the *pds1* mutant which in turn provided the present inventors with important insight for the isolation of a putative cDNA for the *pds1* locus. In animals, genetic defects which inhibit the activity of pOHPP dioxygenase lead to tyrosinemia type I, a fatal inherited disease in aromatic amino acid catabolism characterized by the presence of high levels of pOHPP in the urine.

In an effort to further understand the nature of this disease, pOHPP dioxygenase cDNAs have been cloned from several mammalian and bacterial sources (summarized in Ruetschi et al., Eur. J. Biochem. 205:459-466, 1992). Amino acid identity between various mammalian pOHPP dioxygenase enzymes is >80%; in comparison, their identity to bacterial homologs is very low, less than 28%. By using mammalian and bacterial sequences to search the Expressed Sequence Tags (ESTs) computer DNA database (Newman et al., Plant Physiol. 106:1241-1255, 1994), one partial length *Arabidopsis* EST was identified and used as a probe. The partial length *Arabidopsis* probe corresponds to base pairs 1072 through 1500 of SEQ ID NO:1.

This cDNA contained only 99 amino acids of the carboxyl terminal portion of the protein coding region. The deduced protein sequence of this putative *Arabidopsis* pOHPP dioxygenase cDNA shows similar homology (~50% identity) to both the mammalian and bacterial pOHPP dioxygenases. Interestingly, the partial *Arabidopsis* sequence also contains a 15 amino acid insertion not found in the human or bacterial enzymes. Finally, alignment of six pOHPP dioxygenase sequences from mammals and bacteria identified three regions of high conservation, the highest being a 16 amino acid region near the carboxy end of pOHPP dioxygenases that shows 62.5% identity across all phyla. Ruetschi et al., Eur. J. Biochem. 205:459-466, 1992. This region is also present in the truncated *Arabidopsis* sequence. The lines of evidence suggest that the partial length *Arabidopsis* cDNA described above encodes a pOHPP dioxygenase, most likely the *pds1* locus.

Isolation and Characterization of a full length
Arabidopsis pOHPP dioxygenase cDNA

Utilizing the partial length *Arabidopsis* cDNA probe, an *Arabidopsis* cDNA library was screened by 5 nucleic acid hybridization for full length cDNAs. A large number of hybridizing cDNAs were isolated, and one of the longest, pHPP1.5, containing a 1,520 bp insertion, was sequenced completely; the insert is presented as SEQ ID NO: 1. pHPP1.5 encodes a 446 10 amino acid protein (presented as SEQ ID NO:2), which is slightly larger in size than mammalian and bacterial pOHPP dioxygenases. pHPP1.5 shows 34-40% identity at the amino acid level to pOHPP dioxygenases from various mammals and bacteria. In comparing four 15 bacterial pOHPP dioxygenases and one mammalian pOHPP dioxygenases (pig) which ranged in size from 346-404 amino acids, Denoya et al. identified 69 amino acids that were conserved between all five pOHPP dioxygenases. Denoya et al., J. Bacteriol. 176:5312- 20 5319, 1994. The pHPP1.5 coding region contains 52 of these 69 conserved amino acids.

Demonstration that pHPP1.5 encodes an active pOHPP dioxygenase protein and complements the *pds1* mutation

In order to definitively demonstrate that pHPP1.5 25 is the gene product encoded by the *pds1* locus and that it encodes a functional pOHPP dioxygenase protein, the pHPP1.5 cDNA was cloned into a plant transformation vector for molecular complementation experiments with the *pds1*. The full length wild-type pOHPP dioxygenase 30 cDNA will be subcloned into a plant transformation vector driven by the Cauliflower Mosaic Virus 35S (CaMV) promoter and containing all necessary termination cassettes and selectable markers (Kan^r). The CaMV promoter is a strong constitutive promoter. 35 This single construct and the vector without the pHPP1.5 insert (as a control) will be used in vacuum infiltration transformation which uses whole soil

grown plants and will be done on plants that are heterozygous for the *pds1* mutation. Bouchez et al., CR Acad. Sci. Paris, Sciences de la vie 316, 1993.

In the standard procedure, 20-30 soil grown
5 plants will be independently transformed and analyzed separately. In this case homozygous plants containing the *pds1* mutation would be lethal while heterozygous plants containing the *pds1* mutation would be segregating 2:1 for the *pds1* mutation in their
10 siliques. The inventors will use a similar number of wild type plants in a parallel transformation as a control. After transformation of the *pds1* segregating plant population, as the plants are setting seed the inventors can easily identify those heterozygous for
15 the *pds1* mutation in retrospect by inspection of their siliques which would contain green:white embryos in a 3:1 ratio.

Seed harvested from individually transformed
20 heterozygous *pds1* plants will be germinated on kanamycin and resistant seedlings transferred to soil. Segregation analysis of seed from these primary transformants (T2 seed) and T3 seed for segregation of the *pds1* phenotype (albino and phytoene accumulating) and the T-DNA encoded kanamycin resistance marker
25 (wild type pOHPP dioxygenase cDNA) will conclusively demonstrate complementation of the *pds1* mutation with the pOHPP dioxygenase cDNA. To provide additional proof that pHPP1.5 is encoded by the *pds1* locus, the pHPP1.5 cDNA has been mapped relative to the *pds1*
30 locus using recombinant inbred lines, as described in Lister et al., Plant J. 4:745-750, 1993. The pHPP1.5 cDNA mapped to the region of chromosome 1 containing the *pds1* mutation (Figure 4). Finally, the pHPP1.5 cDNA will be overexpressed in E. coli and the activity
35 of the protein determined.

Modification of pOHPP Expression

From the genetic and biochemical studies described above it is clear that only one pOHPP dioxygenase gene product is involved in chloroplastic quinone synthesis, that the *pds1* mutation defines this gene, that the pHPP1.5 cDNA is the product encoded by the *pds1* locus and that disruption of its function completely eliminates Vitamin E production and plastoquinone and carotenoid synthesis in plant tissues. Modification of pOHPP dioxygenase expression in plants by molecular techniques using pHPP1.5 can therefore be used to positively or negatively affect the production of tocopherols, plastoquinones directly and carotenoids indirectly (refer to Figures 1 and 2). Specifically, overexpression of the pOHPP dioxygenase enzyme will result in increased levels of one or more of these compounds in the tissues of transgenic plants. Alternatively, using antisense techniques, it is possible to lower the level of enzyme activity to decrease the levels of these compounds in plants. Additionally, overexpression of the pOHPP dioxygenase will enable a transgenic plant to withstand elevated levels of herbicides that target this enzyme, providing agronomically significant herbicide resistance relative to normal plants.

Two different plant systems, *Arabidopsis* and tomato, are being used to demonstrate the effects of modified pOHPP dioxygenase in plant tissues. Constitutive overexpression of pOHPP dioxygenase will be done in both plant systems utilizing the CaMV 35S promoter and the pHPP1.5 cDNA. The consequences of this altered expression on tocopherol, plastoquinone and carotenoid levels and profiles in various plant tissues will be determined as described below. In tomato, tissue specific overexpression of pOHPP dioxygenase (pHPP1.5) will be driven by the fruit specific promoter derived from the tomato β subunit.

gene, which is expressed specifically in developing, but not ripening tomato fruit. This will determine the potential for modifying the levels of tocopherol, plastoquinone and carotenoids specifically in
5 developing and ripening fruit for nutritional purposes without affecting their production in other plant tissues. These combined experiments will determine whether pOHPP dioxygenase is a rate limiting step in chloroplastic homogentisic acid derived quinone
10 synthesis and the potential for manipulating chloroplastic homogentisic acid derived quinones (tocopherols and plastoquinones) and compounds that require quinones for their synthesis (carotenoids, etc) by increasing pOHPP dioxygenase activity.
15 Multiple independent transformants will be produced for each construct and plant species used. The integration and gene copy number of each chimeric gene in each line will be confirmed by southern analysis, the level of pOHPP mRNA determined by
20 Northern blot analysis, pOHPP dioxygenase activity determined as described in Schulz et al., FEBS 318:162-166, 1993, and the effects on individual chloroplastic components of interest analyzed (tocopherols, plastoquinones and carotenoids). In
25 green tissue containing constitutively expressing constructs this analysis can occur relatively soon after transformants are put into soil. Analysis of fruit specific construct lines will require much more time for fruit set to occur. Analysis of tocopherols, plastoquinones and carotenoids will be by a combination of HPLC, optical and mass spectra as described in Norris et al. (1995, in press). Analysis of tocopherol levels is performed by HPLC and when needed by GC:mass spectroscopy in selected ion mode.
30 In MS analysis the absolute level of tocopherol will be quantified by isotopic dilution with a known, "heavy carbon" tocopherol standard added at the start
35

of the extraction. Determination based on fresh weight of tissue can also be performed. Plastoquinone levels will be quantified by C8 HPLC and optical spectra as described in Norris et al. (1995, in press). Total carotenoid levels are determined spectrophotometrically and the levels of individual carotenes quantified by C18 HPLC and optical spectra quantified to standards. In the course of these experiments we will identify high expressing lines with simple insertions that segregate as single genetic loci in progeny. This will facilitate analysis of the inheritance of the gene and phenotype in future generations.

Overexpression of pOHPP for in vitro Herbicide Analysis

pOHPP dioxygenase will be overexpressed in E. coli or other prokaryotic or eukaryotic protein production systems and purified in large amounts for use in enzymatic assays for identifying new herbicide compounds (pOHPP inhibitors) and optimizing existing chemistries through detailed kinetic analysis.

SEQUENCE LISTING

(1) GENERAL INFORMATION:

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5 (ii) TITLE OF INVENTION: Cloned Plant P-Hydroxyphenyl
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(B) COMPUTER: IBM PC compatible
(C) OPERATING SYSTEM: PC-DOS/MS-DOS
(D) SOFTWARE: PatentIn Release #1.0, Version #1.30

25 (vi) CURRENT APPLICATION DATA:
(A) APPLICATION NUMBER:
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(2) INFORMATION FOR SEQ ID NO:1:

35 (i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 1519 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

40 (vi) ORIGINAL SOURCE:
(A) ORGANISM: Arabidopsis thaliana

(vii) IMMEDIATE SOURCE:
(B) CLONE: pHPP1.5

(ix) FEATURE:
(A) NAME/KEY: CDS
45 (B) LOCATION: 37..1374

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

	CCACGCGTCC GAGTTTAGC AGAGTTGGTG AAATCA ATG GGC CAC CAA AAC GCC Met Gly His Gln Asn Ala	54
5	GCC GTT TCA GAG AAT CAA AAC CAT GAT GAC GGC GCT GCG TCG TCG CCG Ala Val Ser Glu Asn Gln Asn His Asp Asp Gly Ala Ala Ser Ser Pro 10 15 20	102
10	GGA TTC AAG CTC GTC GGA TTT TCC AAG TTC GTA AGA AAG AAT CCA AAG Gly Phe Lys Leu Val Gly Phe Ser Lys Phe Val Arg Lys Asn Pro Lys 25 30 35	150
15	TCT GAT AAA TTC AAG GTT AAG CGC TTC CAT CAC ATC GAG TTC TGG TGC Ser Asp Lys Phe Lys Val Lys Arg Phe His His Ile Glu Phe Trp Cys 40 45 50	198
20	GGC GAC GCA ACC AAC GTC GCT CGT CGC TTC TCC TGG GGT CTG GGG ATG Gly Asp Ala Thr Asn Val Ala Arg Arg Phe Ser Trp Gly Leu Gly Met 55 60 65 70	246
25	AGA TTC TCC GCC AAA TCC GAT CTT TCC ACC GGA AAC ATG GTT CAC GCC Arg Phe Ser Ala Lys Ser Asp Leu Ser Thr Gly Asn Met Val His Ala 75 80 85	294
30	TCT TAC CTA CTC ACC TCC GGT GAC CTC CGA TTC CTT TTC ACT GCT CCT Ser Tyr Leu Leu Thr Ser Gly Asp Leu Arg Phe Leu Phe Thr Ala Pro 90 95 100	342
35	TAC TCT CCG TCT CTC TCC GCC GGA GAG ATT AAA CCG ACA ACC ACA GCT Tyr Ser Pro Ser Leu Ser Ala Gly Glu Ile Lys Pro Thr Thr Thr Ala 105 110 115	390
40	TCT ATC CCA AGT TTC GAT CAC GGC TCT TGT CGT TCC TTC TTC TCT TCA Ser Ile Pro Ser Phe Asp His Gly Ser Cys Arg Ser Phe Phe Ser Ser 120 125 130	438
45	CAT GGT CTC GGT GTT AGA GCC GTT GCG ATT GAA GTA GAA GAC GCA GAG His Gly Leu Gly Val Arg Ala Val Ala Ile Glu Val Glu Asp Ala Glu 135 140 145 150	486
50	TCA GCT TTC TCC ATC AGT GTA GCT AAT GGC GCT ATT CCT TCG TCG CCT Ser Ala Phe Ser Ile Ser Val Ala Asn Gly Ala Ile Pro Ser Ser Pro 155 160 165	534
55	CCT ATC GTC CTC AAT GAA GCA GTT ACG ATC GCT GAG GTT AAA CTA TAC Pro Ile Val Leu Asn Glu Ala Val Thr Ile Ala Glu Val Lys Leu Tyr 170 175 180	582
60	GGC GAT GTT GTT CTC CGA TAT GTT AGT TAC AAA GCA GAA GAT ACC GAA Gly Asp Val Val Leu Arg Tyr Val Ser Tyr Lys Ala Glu Asp Thr Glu 185 190 195	630
65	AAA TCC GAA TTC TTG CCA GGG TTC GAG CGT GTA GAG GAT GCG TCG TCG Lys Ser Glu Phe Leu Pro Gly Phe Glu Arg Val Glu Asp Ala Ser Ser 200 205 210	678
70	TTC CCA TTG GAT TAT GGT ATC CGG CGG CTT GAC CAC GCC GTG GGA AAC Phe Pro Leu Asp Tyr Gly Ile Arg Arg Leu Asp His Ala Val Gly Asn 215 220 225 230	726
75	GTT CCT GAG CTT GGT CCG GCT TTA ACT TAT GTA GCG GGG TTC ACT GGT Val Pro Glu Leu Gly Pro Ala Leu Thr Tyr Val Ala Gly Phe Thr Gly 235 240 245	774

	TTT CAC CAA TTC GCA GAG TTC ACA GCA GAC GAC GTT GGA ACC ACC GCC GAG Phe His Gln Phe Ala Glu Phe Thr Ala Asp Asp Val Gly Thr Ala Glu 250 255 260	822
5	AGC GGT TTA AAT TCA GCG GTC CTG GCT AGC AAT GAT GAA ATG GTT CTT Ser Gly Leu Asn Ser Ala Val Leu Ala Ser Asn Asp Glu Met Val Leu 265 270 275	870
	CTA CCG ATT AAC GAG CCA GTG CAC GGA ACA AAG AGG AAG AGT CAG ATT Leu Pro Ile Asn Glu Pro Val His Gly Thr Lys Arg Lys Ser Gln Ile 280 285 290	918
10	CAG ACG TAT TTG GAA CAT AAC GAA GGC GCA GGG CTA CAA CAT CTG GCT Gln Thr Tyr Leu Glu His Asn Glu Gly Ala Gly Leu Gln His Leu Ala 295 300 305 310	966
15	CTG ATG AGT GAA GAC ATA TTC AGG ACC CTG AGA GAG ATG AGG AAG AGG Leu Met Ser Glu Asp Ile Phe Arg Thr Leu Arg Glu Met Arg Lys Arg 315 320 325	1014
	AGC AGT ATT GGA GGA TTC GAC TTC ATG CCT TCT CCT CCG CCT ACT TAC Ser Ser Ile Gly Gly Phe Asp Phe Met Pro Ser Pro Pro Pro Thr Tyr 330 335 340	1062
20	TAC CAG AAT CTC AAG AAA CGG GTC GGC GAC GTG CTC AGC GAT GAT CAG Tyr Gln Asn Leu Lys Arg Val Gly Asp Val Leu Ser Asp Asp Gln 345 350 355	1110
	ATC AAG GAG TGT GAG GAA TTA GGG ATT CTT GTA GAC AGA GAT GAT CAA Ile Lys Glu Cys Glu Leu Gly Ile Leu Val Asp Arg Asp Asp Gln 360 365 370	1158
25	GGG ACG TTG CTT CAA ATC TTC ACA AAA CCA CTA GGT GAC AGG CCG ACG Gly Thr Leu Leu Gln Ile Phe Thr Lys Pro Leu Gly Asp Arg Pro Thr 375 380 385 390	1206
30	ATA TTT ATA GAG ATA ATC CAG AGA GTA GGA TGC ATG ATG AAA GAT GAG Ile Phe Ile Glu Ile Ile Gln Arg Val Gly Cys Met Met Lys Asp Glu 395 400 405	1254
	GAA GGG AAG GCT TAC CAG AGT GGA GGA TGT GGT GGT TTT GGC AAA GGC Glu Gly Lys Ala Tyr Gln Ser Gly Gly Cys Gly Gly Phe Gly Lys Gly 410 415 420	1302
35	AAT TTC TCT GAG CTC TTC AAG TCC ATT GAA GAA TAC GAA AAG ACT CTT Asn Phe Ser Glu Leu Phe Lys Ser Ile Glu Glu Tyr Glu Lys Thr Leu 425 430 435	1350
	GAA GCC AAA CAG TTA GTG GGA TGA ACAAGAAGAA GAACCAACTA AAGGATTGTG Glu Ala Lys Gln Leu Val Gly * 440 445	1404
40	TAATTAATGT AAAACTGTT TATCTTATCA AAACAATGTT ATACAACATC TCATTTAAAAA ACGAGATCAA TCAAAAAATA CAATCTTAAA TTCAAAACCA AAAAAAAA AAAAAA (2) INFORMATION FOR SEQ ID NO:2:	1464 1519

- 45 (i) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 446 amino acids.
 (B) TYPE: amino acid
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Met Gly His Gln Asn Ala Ala Val Ser Glu Asn Gln Asn His Asp Asp
 1 5 10 15

Gly Ala Ala Ser Ser Pro Gly Phe Lys Leu Val Gly Phe Ser Lys Phe
 5 20 25 30

Val Arg Lys Asn Pro Lys Ser Asp Lys Phe Lys Val Lys Arg Phe His
 35 40 45

His Ile Glu Phe Trp Cys Gly Asp Ala Thr Asn Val Ala Arg Arg Phe
 50 55 60

Ser Trp Gly Leu Gly Met Arg Phe Ser Ala Lys Ser Asp Leu Ser Thr
 10 65 70 75 80

Gly Asn Met Val His Ala Ser Tyr Leu Leu Thr Ser Gly Asp Leu Arg
 85 90 95

Phe Leu Phe Thr Ala Pro Tyr Ser Pro Ser Leu Ser Ala Gly Glu Ile
 15 100 105 110

Lys Pro Thr Thr Ala Ser Ile Pro Ser Phe Asp His Gly Ser Cys
 115 120 125

Arg Ser Phe Phe Ser Ser His Gly Leu Gly Val Arg Ala Val Ala Ile
 130 135 140

Glu Val Glu Asp Ala Glu Ser Ala Phe Ser Ile Ser Val Ala Asn Gly
 20 145 150 155 160

Ala Ile Pro Ser Ser Pro Pro Ile Val Leu Asn Glu Ala Val Thr Ile
 165 170 175

Ala Glu Val Lys Leu Tyr Gly Asp Val Val Leu Arg Tyr Val Ser Tyr
 25 180 185 190

Lys Ala Glu Asp Thr Glu Lys Ser Glu Phe Leu Pro Gly Phe Glu Arg
 195 200 205

Val Glu Asp Ala Ser Ser Phe Pro Leu Asp Tyr Gly Ile Arg Arg Leu
 210 215 220

Asp His Ala Val Gly Asn Val Pro Glu Leu Gly Pro Ala Leu Thr Tyr
 30 225 230 235 240

Val Ala Gly Phe Thr Gly Phe His Gln Phe Ala Glu Phe Thr Ala Asp
 245 250 255

Asp Val Gly Thr Ala Glu Ser Gly Leu Asn Ser Ala Val Leu Ala Ser
 35 260 265 270

Asn Asp Glu Met Val Leu Leu Pro Ile Asn Glu Pro Val His Gly Thr
 275 280 285

Lys Arg Lys Ser Gln Ile Gln Thr Tyr Leu Glu His Asn Glu Gly Ala
 290 295 300

Gly Leu Gln His Leu Ala Leu Met Ser Glu Asp Ile Phe Arg Thr Leu
 40 305 310 315 320

Arg Glu Met Arg Lys Arg Ser Ser Ile Gly Gly Phe Asp Phe Met Pro
 325 330 335

Ser Pro Pro Pro Thr Tyr Tyr Gln Asn Leu Lys Lys Arg Val Gly Asp
 45 340 345 350

Val Leu Ser Asp Asp Gln Ile Lys Glu Cys Glu Glu Leu Gly Ile Leu
355 360 365

Val Asp Arg Asp Asp Gln Gly Thr Leu Leu Gln Ile Phe Thr Lys Pro
370 375 380

5 Leu Gly Asp Arg Pro Thr Ile Phe Ile Glu Ile Ile Gln Arg Val Gly
385 390 395 400

Cys Met Met Lys Asp Glu Glu Gly Lys Ala Tyr Gln Ser Gly Gly Cys
405 410 415

10 Gly Gly Phe Gly Lys Gly Asn Phe Ser Glu Leu Phe Lys Ser Ile Glu
420 425 430

Glu Tyr Glu Lys Thr Leu Glu Ala Lys Gln Leu Val Gly *
435 440 445

5

CLAIMS

I claim:

1. A biologically pure sample of DNA, the DNA comprising a DNA sequence coding for the expression of a plant p-hydroxyphenyl pyruvic acid dioxygenase.
- 10 2. A vector containing the DNA sequence of claim 1.
3. A microbial host transformed by the vector of claim 2.
- 15 4. The DNA of claim 1, wherein the p-hydroxyphenyl pyruvic acid dioxygenase is from *Arabidopsis thaliana*.
5. A transgenic tomato plant transformed with a DNA construct including the DNA of claim 1.
- 20 6. A transgenic *Arabidopsis* plant transformed with a DNA construct including the DNA of claim 1.
7. The biologically pure DNA of claim 1 wherein the DNA is SEQ ID NO:2.
- 25 8. A DNA plant gene expression construct comprising:
 - a. a DNA sequence coding for the expression of a plant p-hydroxyphenyl pyruvic acid dioxygenase;
 - b. a promoter effective in plant cells located 5' to the DNA coding sequence; and
 - 30 c. a 3' termination sequence effective in plant cells.

9. A DNA construct comprising:

- a. a promoter capable of expressing a downstream coding sequence in a tomato plant;
- b. a DNA sequence coding for the expression of a p-hydroxyphenyl pyruvic acid dioxygenase of plant origin; and
- c. a 3' termination sequence, the construction capable of expressing a p-hydroxyphenyl pyruvic acid dioxygenase gene when transformed into tomato plants.

10. A bacteria containing the construction of Claim 9.

11. A tomato plant cell containing the construction of Claim 9.

15 12. An Arabidopsis plant cell containing the construction of Claim 9.

20 13. A transgenic tomato plant comprising in its genome a foreign genetic construction comprising, 5' to 3', a promoter effective in tomato, a DNA coding region encoding p-hydroxyphenyl pyruvic acid dioxygenase, and a transcriptional terminator, the genetic construction effective in vivo in tomato plants to stimulate expression of p-hydroxyphenyl pyruvic acid dioxygenase.

25 14. Seed of the tomato plant of claim 13.

15. Fruit of the tomato plant of claim 13.

30 16. The transgenic tomato plant of claim 11 wherein the DNA coding region encoding p-hydroxyphenyl pyruvic acid dioxygenase is that set forth in SEQ ID NO:1.

17. A method of suppressing the production of vitamin E and plastoquinones in a plant, the method comprising the steps of:
- a. isolating a DNA sequence encoding a p-hydroxyphenyl pyruvic acid dioxygenase of plant origin;
 - b. creating a genetic construction including, 5' to 3', a promoter effective in the plant's cells, a coding sequence, and a transcriptional terminator, the coding region being derived from the DNA sequence, wherein the DNA sequence from step (a) has been altered so that expression of p-hydroxyphenyl pyruvic acid dioxygenase is suppressed; and
 - c. transforming a cell of the plant with the 10 genetic construction, whereby the plant's cell produces lowered levels of vitamin E and plastoquinones.

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Carotenoid Pathway

Tocopherol/Plastoquinone Pathway

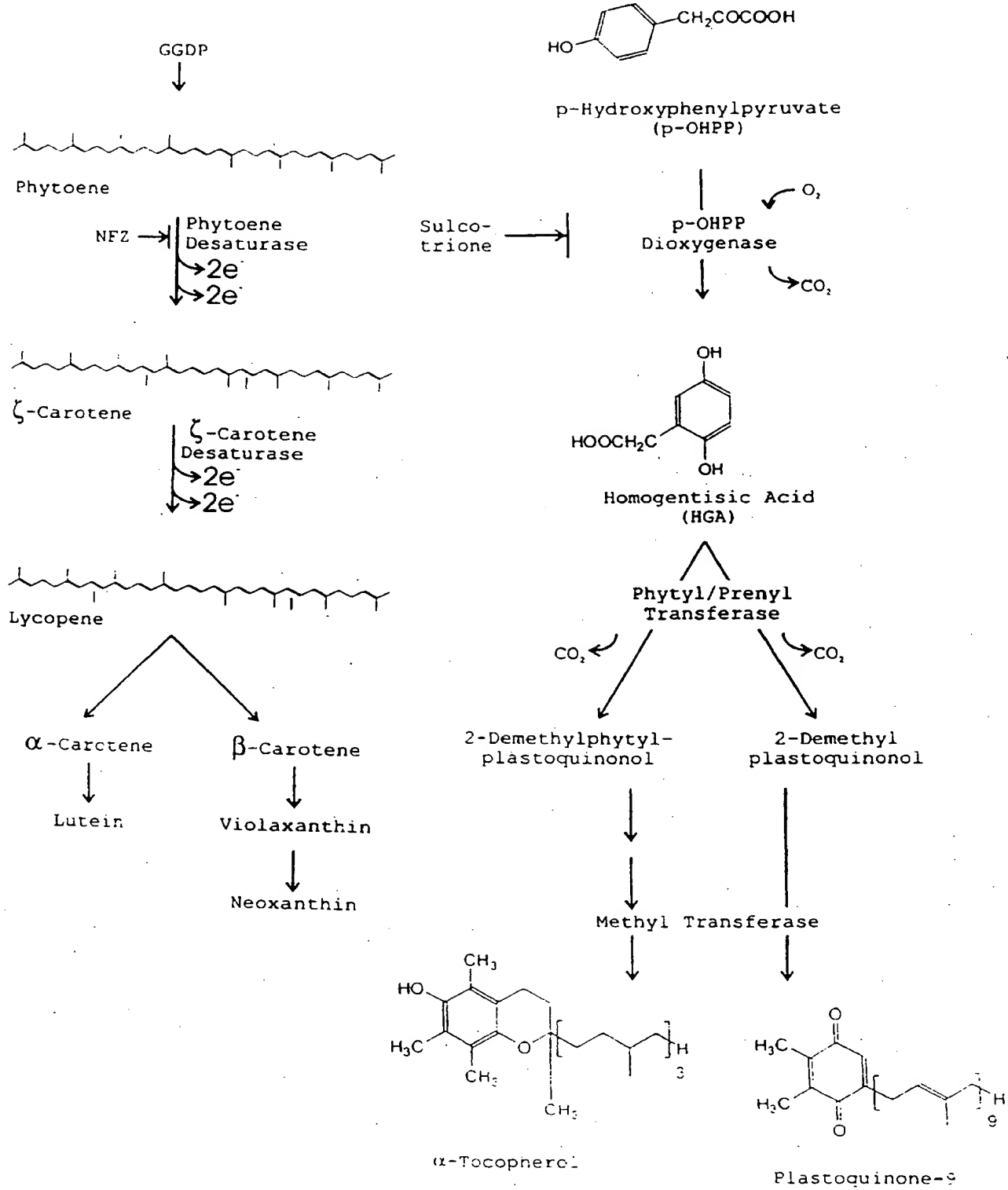


FIG 1

SUBSTITUTE SHEET (RULE 26)

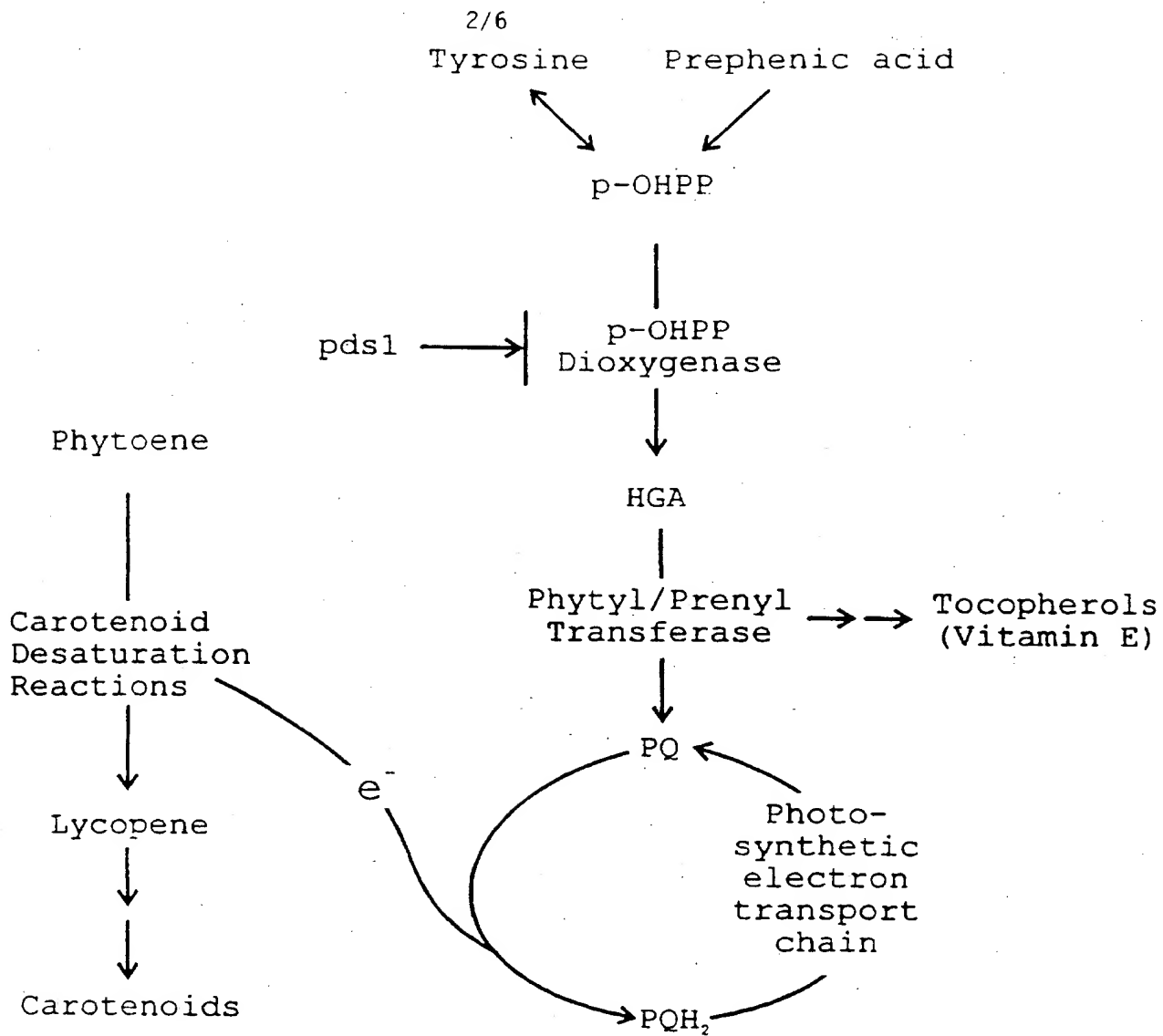


FIG 2

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FIG 3A

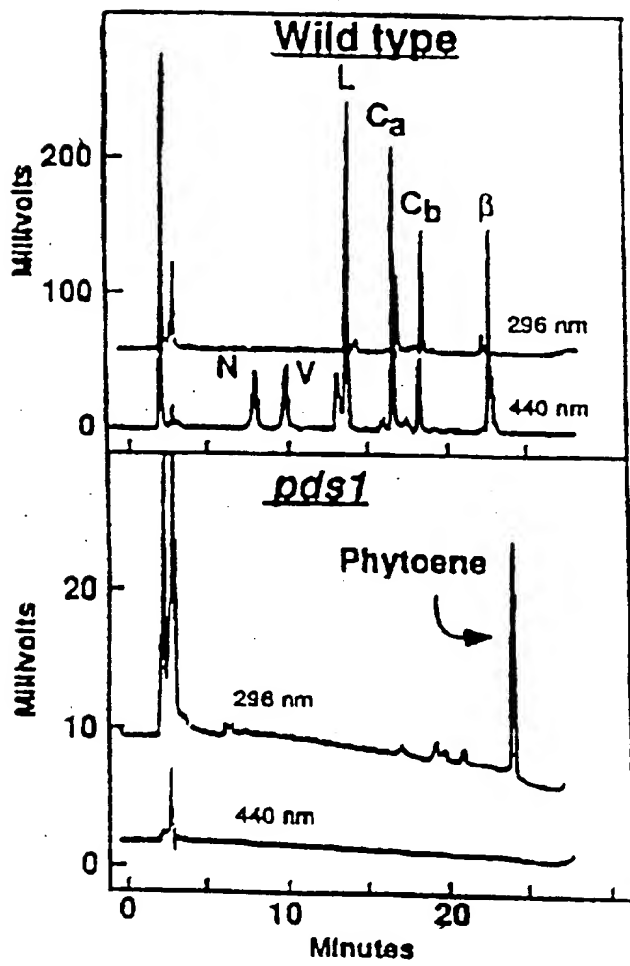


FIG 3B

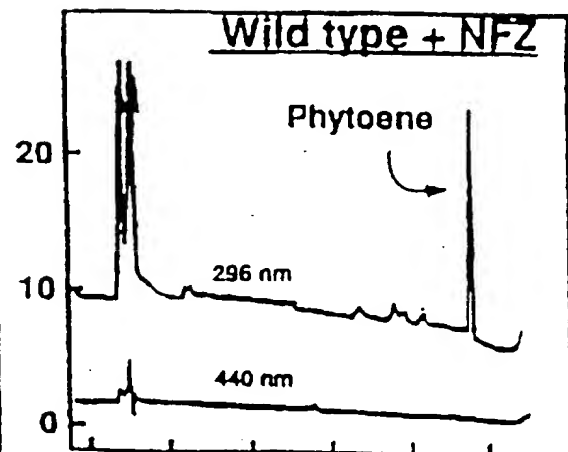


FIG 3D

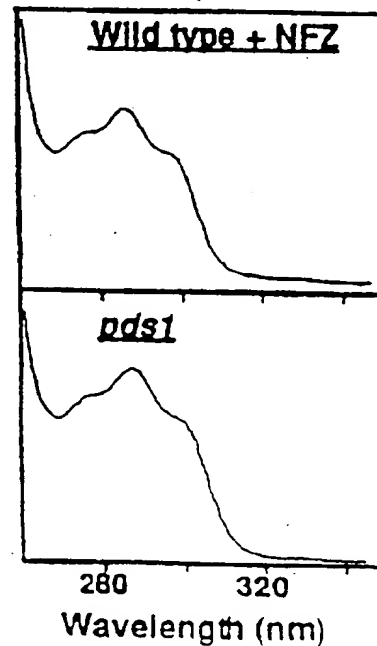


FIG 3E

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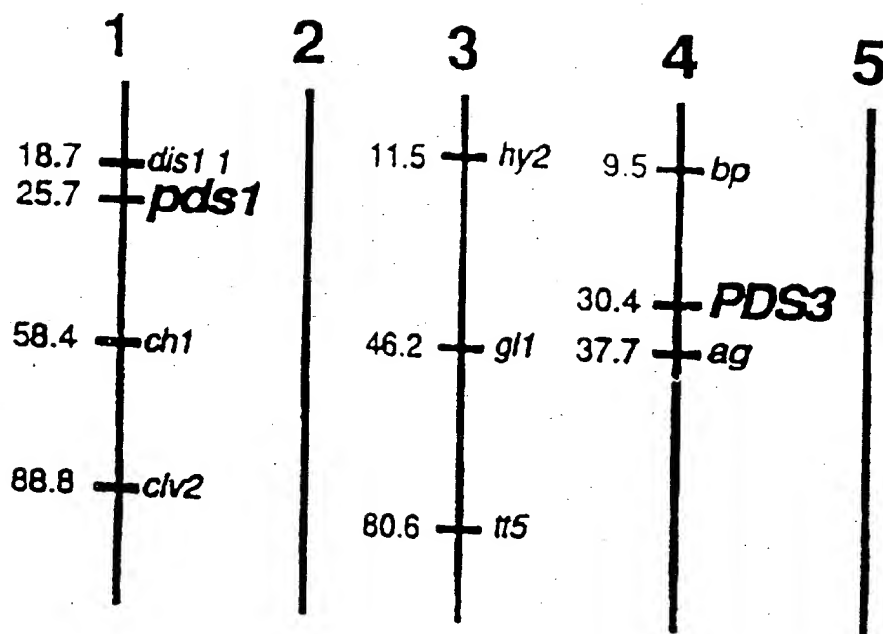
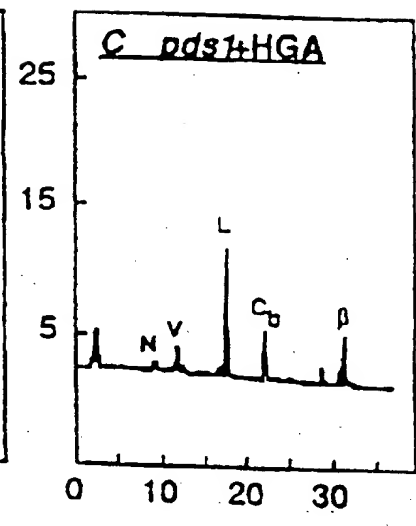
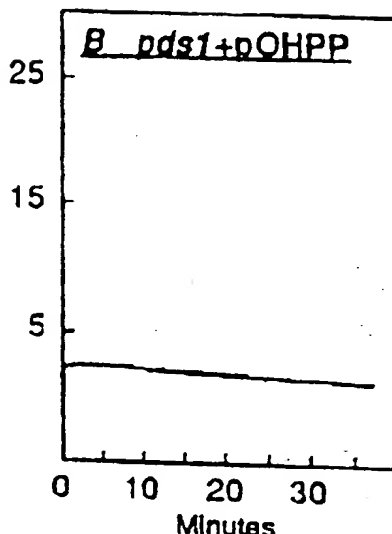
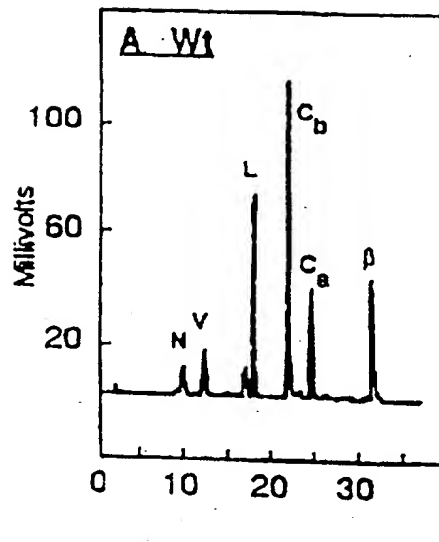


FIG 4

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**FIG 5A****FIG 5B****FIG 5C**

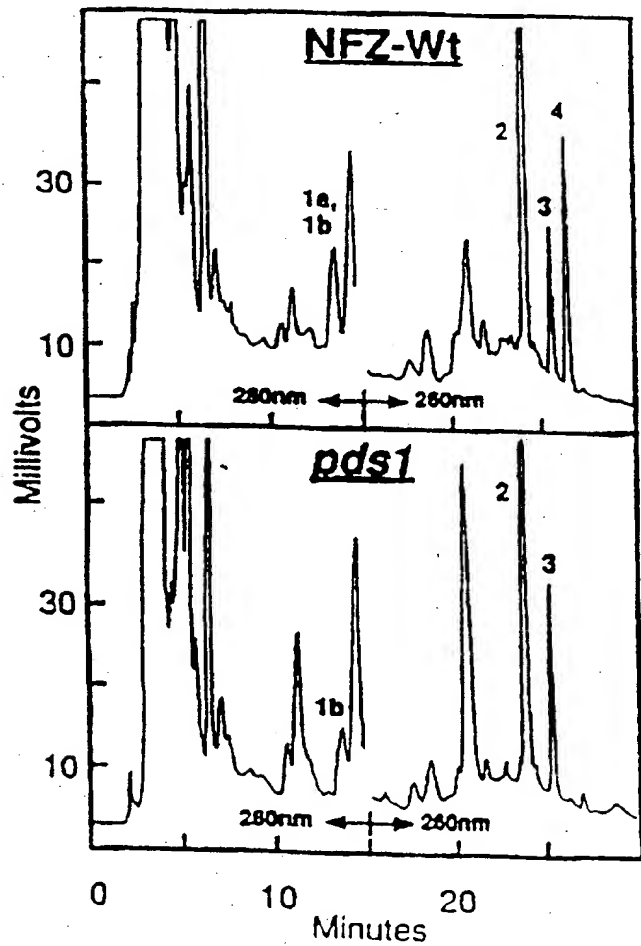
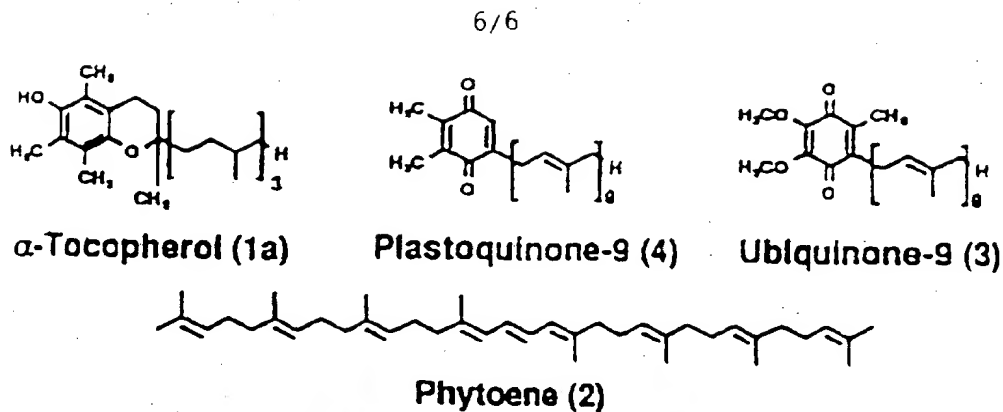


FIG 6A

FIG 6B